(19) World Intellectual Property Organization International Bureau



(43) International Publication Date 8 March 2001 (08.03.2001)

PCT

(10) International Publication Number WO 01/16285 A2

(51) International Patent Classification7:

C12N

- (21) International Application Number: PCI/DK00/00476
- (22) International Filing Date: 31 August 2000 (31.08.2000)
- (25) Filing Language:

English

(26) Publication Language:

English

(30) Priority Data:

- PA 1999 01212 31 August 1999 (31.08.1999) DK PA 1999 01500 20 October 1999 (20.10.1999) DK
- (71) Applicant: NOVOZYMES A/S [DK/DK]; Krogshoejvej 36, DK-2880 Bagsvaerd (DK).
- (72) Inventors: NØRREGAARD-MADSEN, Mads; Munkerisvej 31 2tv, DK-5230 Odense M (DK). RAHBEK ØSTERGAARD, Peter; Kvædevej 111, DK-2830 Virum (DK). VØGE CHRISTENSEN, Claus, Bo; G. A Hagemannsvej 13, DK-3070 Snekkersten (DK). FLENSTED LASSEN, Søren; Lyngbyvej 19, lejl. 17, DK-2100 København Ø (DK).

- (81) Designated States (national): AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN, CR, CU, CZ, DE, DK, DM, DZ, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, UZ, VN, YU, ZA, ZW.
- (84) Designated States (regional): ARIPO patent (GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZW), Eurasian patent (AM, AZ, BY, KG, KZ, MD, RU, TJ, TM), European patent (AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE), OAPI patent (BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG).

Published:

 Without international search report and to be republished upon receipt of that report.

For two-letter codes and other abbreviations, refer to the "Guidance Notes on Codes and Abbreviations" appearing at the beginning of each regular issue of the PCT Gazette.

5 A

(54) Title: NOVEL PROTEASES AND VARIANTS THEREOF

(57) Abstract: Novel isolated proteases of the RP-II type and variants of RP-II proteases exhibiting improved properties in comparison to the parent RP-II protease, DNA constructs and vectors coding for the expression of said proteases and variants, host cells capable of expressing the proteases and variants from the DNA constructs, as well as a method of producing them by cultivating said host cells. The proteases may advantageously be used as constituents in detergent compositions and additives, optionally in combination with other enzymes such as proteases, lipases, cellulases, amylases, peroxidases or oxidases.

5

10

20

25

30

35

40

NOVEL PROTEASES AND VARIANTS THEREOF

FIELD OF THE INVENTION

The present invention relates to novel isolated proteases of the RP-II type and variants of RP-II proteases exhibiting improved properties in comparison to the parent RP-II protease, DNA constructs and vectors coding for the expression of said proteases and variants, host cells capable of expressing the proteases and variants from the DNA constructs, as well as a method of producing them by cultivating said host cells. The proteases may advantageously be used as constituents in detergent compositions and additives.

BACKGROUND OF THE INVENTION

Protein Science 6 (1997) 501-523) have for many years been used in the detergent industry due to their apparent superiority over other protease types in this type of application.

A large number of subtilisins and the related subtilases are known.

Protease variants have been produced in a number of subtilisin proteases in order to provide changes in various properties, such as thermo stability, specific activity, pH-dependency, isoelectric point, wash performance, oxidation stability, autoproteolysis, etc.

Such variants are disclosed in various patent publications, such as EP 130 756, EP 251 446, and EP 824 585.

The fact that novel detergents constantly are being developed to satisfy various user demands provides an incentive to continuously develop novel proteases capable of providing excellent performance in the novel detergents.

Bacillus proteases of the RP-II type are another type of serine proteases that in primary structure are similar to chymotrypsinogen.

The first description of a protease of the RP-II family of Bacillus proteases was in US Patent No. 4,266,031 (Tang et al., Novo Industri A/S), where it was designated Component C and tentatively characterised as not being a serine protease or metallo protease. Component C was considered a contaminant

in the production of the *Bacillus licheniformis* alkaline protease, subtilisin Carlsberg.

In EP 369 817 (Omnigene Bioproducts, Inc.) the B. subtilis member of the RP-II family was identified by its amino acid and DNA sequences. The enzyme was stated not to be a serine protease, and the family name RP-II designated (Residual Protease II). The enzyme was characterised further as a metallo protease by the inventors of EP 369 817 (Rufo et al., 1990, J. Bacteriol. 2 1019-1023, and Sloma et al., 1990, J. Bacteriol. 2 1024-1029), designating the enzyme as mpr.

In WO 91/13553 (Novo Nordisk A/S) the amino acid sequence of the C component was disclosed, stating that it is a serine protease specific for glutamic and aspartic acid, while EP 482 879 (Shionogi & Co. Ltd.) disclosed the enzyme and a DNA sequence encoding the C component from B. licheniformis ATCC No. 14580, naming the enzyme BLase. In EP 482 879 the protease is described as being specific for glutamic acid.

In 1997 Okamoto et al. (Appl. Microbiol. Biotechnol. (1997) 48 27-33) found that the B. subtilis homologue of BLase, named BSase was identical to the above-mentioned enzyme, mpr/RP-II.

SUMMARY OF THE INVENTION

10

15

20

25

30

35

Initial testing of the *B. licheniformis* member of the RP-II family indicated that this enzyme in some aspects might be inferior in detergents in comparison to the subtilisins.

However, it is believed that a screening program for novel RP-II family members both isolated from nature (wild types) and recombinantly produced variants thereof will provide alternative proteases for use in detergents.

Consequently it is an object of the present invention to provide novel RP-II protease members obtainable from various Bacillus strains.

Furthermore it is the object of the present invention to design novel variants of the RP-II proteases having improved properties as compared to those of their parent protease.

Accordingly, in a first aspect the present invention relates to novel isolated RP-II proteases selected from the group consisting of:

(i) a RP-II protease that is immunochemically identical or partially identical by cross-reaction with an antibody raised against or reactive with at least one epitope of a RP-II protease comprising the amino acid sequences of the mature peptides shown in the appended Sequence Listing SEQ ID NO: 2, 4, ID No. 6, ID No. 8, ID No. 10, or ID No. 12; and/or

5

25

35

40

- 10 (ii) a RP-II protease that is at least 60% homologous with the amino acid sequence of a RP-II protease comprising the amino acid sequence shown in the appended Sequence Listing SEQ ID NO: 2, SEQ ID NO: 4, SEQ ID NO: 6, SEQ ID NO: 8, SEQ ID NO: 10, or SEQ ID NO: 12; and/or
- 15 (iiia) a RP-II protease that is encoded by a DNA sequence which hybridizes with an oligonucleotide probe hybridizing with a DNA sequence encoding a RP-II protease comprising the amino acid sequence shown in the appended Sequence Listing SEQ ID NO: 2, SEQ ID NO: 4, SEQ ID NO: 6, SEQ ID NO: 20 8, SEQ ID NO: 10, or SEQ ID NO: 12; and/or
 - (iiib) a RP-II protease that is encoded by a DNA sequence which hybridizes with an oligonucleotide probe hybridizing with a DNA sequence encoding a RP-II protease comprising the DNA sequence shown in the appended Sequence Listing SEQ ID NO: 1, SEQ ID NO: 3, SEQ ID NO: 5, SEQ ID NO: 7, SEQ ID NO: 9, or SEQ ID NO: 11;
 - (iv) an allelic variant of (i), (ii), or (iiia or
 iiib);
- (v) a subsequence of (i), (ii), (iiia or iiib), or 30 (iv), wherein the subsequence has protease activity.

The invention furthermore relates to RP-II protease variants produced by modifying at least one amino acid residue within the mature enzyme in order to obtain a modification of the properties of the parent enzyme.

Such variants of the present invention are contemplated to have improved substrate specificities, catalytic rate, stability, especially towards the action of proteolytic enzymes and/or detergent ingredients, thermostability, storage stability, improved resistance towards peroxidase/pHBS inactivation, and/or improved wash performance as compared to

the parent RP-II protease. The variants of the invention include fragments of the RP-II proteases or variants thereof having retained protease activity.

The present invention also relates to novel isolated nucleic acid sequences encoding RP-II proteases, selected from the group consisting of:

- (a) a nucleic acid sequence having at least 60% homology with the nucleic acid sequence encoding the mature polypeptide of SEQ ID NO: 1, 3, 5, 7, 9, or 11;
- 10 (b) a nucleic acid sequence which hybridizes under low stringency conditions with (i) the nucleic acid sequence of SEQ ID NO: 1, 3, 5, 7, 9, or 11 (ii) the cDNA sequence of SEQ ID NO: 1, 3, 5, 7, 9, or 11, (iii) a subsequence of (i) or (ii) of at least 100 nucleotides, or (iv) a complementary strand of (i), (ii), or (iii);
 - (c) an allelic variant of (a), or (b);
 - (d) a subsequence of (a), (b), or (c), wherein the subsequence encodes a polypeptide fragment which has protease activity; and

The present invention also relates to nucleic acid or DNA constructs comprising a DNA sequence encoding a RP-II protease or RP-II protease variant as indicated above, recombinant expression vectors carrying said DNA construct, cells transformed with a DNA construct or expression vector, as well as methods for producing a RP-II protease or variant of the invention by culturing or growing said cell under conditions conducive to the production of the protease or variant, after which the protease or variant is recovered from the culture, and optionally purified to be substantially pure.

The invention further relates to an enzyme granulate, a liquid enzyme composition or a protected enzyme preparation comprising a RP-II protease or protease variant of the invention and suitable for the preparation of e.g. a detergent composition comprising a RP-II protease or RP-II protease variant of the invention.

DEPOSITED BIOLOGICAL MATERIALS

20

25

30

35

40

DNA encoding the novel RP-II proteases of the invention has been inserted into plasmids used to transform *E. coli*. These transformants have been deposited according to the

Budapest Treaty on the International Recognition of the Deposits of Microorganisms for the Purpose of Patent Procedures, on 3 December 1990 at DSMZ-Deutsche Sammlung von Mikroorganismen und Zellkulturen GmbH (DSM), Mascheroder Weg 1b, D-38124 Braunschweig, Germany, under Accession Nos. DSM 12841 (AC116), DSM 12842 (CDJ31), DSM 12843 (BO32), DSM 12844 (JA96), and DSM 12845 (AA519).

The deposits have been made under conditions that assure that access to the culture will be available during the pendency of this patent application to one determined by the Commissioner of Patents and Trademarks to be entitled thereto under 37 C.F.R. §1.14 and 35 U.S.C. §122. The deposit represents a substantially pure culture of the deposited strain. The deposit is available as required by foreign patent laws in countries wherein counterparts of the subject application, or its progeny are filed. However, it should be understood that the availability of a deposit does not constitute a license to practice the subject invention in derogation of patent rights granted by governmental action.

20

25

35

10

15

BRIEF DESCRIPTION OF DRAWINGS

The present invention is further illustrated by reference to the accompanying drawings, in which:

Figs. 1a to 1c shows an alignment of the novel wild type RP-II proteases to the RP-II protease from Bacillus licheniformis, BCL, in the manner described below to establish the numbering of the amino acid residues for each novel wild type protease.

Figs. 2a and 2b shows schematically the construction of 30 plasmid pNM1003.

DEFINITIONS

In the present context, the term "RP-II protease" is intended to indicate an evolutionary homologue of the RP-II protease derived from a bacterium of the genus Bacillus, and in particular of any of the species B. licheniformis, B. pumilus, B. subtilis, or B. halmapalus or a functional analogue thereof.

The term "functional analogue" is intended to indicate 40 a RP-II protease which is immunologically cross-reactive with

at least one of the RP-II proteases described herein, and/or comprises an amino acid sequence which is more than 60% homologous with that of at least one of the mature RP-II proteases shown in SEQ ID NO: 2, SEQ ID NO: 4, SEQ ID NO: 6, SEQ ID NO: 8, SEQ ID NO: 10, or SEQ ID NO: 12, such as more than 70%, 80% or even 90% homologous with said proteases, is encoded by a DNA sequence hybridizing with an oligonucleotide probe which also hybridizes with at least one of the DNA sequences encoding the RP-II proteases, the DNA and amino acid sequences of which are shown in SEQ ID NO: 1, SEQ ID NO: 3, SEQ ID NO: 5, SEQ ID NO: 7, SEQ ID NO: 9, and SEQ ID NO: 11.

10

15

20

25

30

35

In the context of this application the term "homologue" or "homologous" is meant to comprise other parent (wild-type) RP-II proteases, which have a primary structure similar to that of another RP-II protease. The homology between two amino acid sequences is in this context described by the parameter "identity".

Sequence comparisons can be performed by standard methods, such as the Wilbur-Lipman method (Wilbur and Lipman, 1983, Proceedings of the National Academy of Science USA 80: 726-730), the Clustal method (Higgins, 1989, CABIOS 5: 151-153), or the GCG method Needleman, S.B. and Wunsch, C.D., (1970), Journal of Molecular Biology, 48, 443-453)

In order to determine the degree of identity between two RP-II proteases the GAP routine of the GCG package version 9.1 (Genetics Computer Group, 575 Science Drive, Madison, Wisconsin, USA 53711) can be applied using the following parameters: gap creation penalty = 8 and gap extension penalty = 8 and all other parameters kept at their default values. The output from the routine is besides the calculation of the "Percent Identity" between the two sequences the amino acid alignment between the two sequences.

Based on this it is routine for a person skilled in the art to identify suitable homologous RP-II proteases and corresponding homologous positions, which can be modified according to the invention.

The term "parent" as used herein is typically a wild type protease, meaning that it has been described or found to be produced by a Bacillus species or strain isolated from 40 natural sources. However, parent is also meant to comprise any

٠ جيتيت

protease, such as a variant protease, being used as starting material for further modifications to produce a further variant.

intended "variant" is to indicate The term polypeptide which is derived from a RP-II protease as defined above and which has one or more of the properties i)-iii) which will be further discussed below. Typically, the variant differ from the RP-II protease by one or more amino acid residues, which, for instance, may have been added or deleted from either or both of the N-terminal or C-terminal end of the protease, inserted or deleted at one or more sites within the amino acid sequence of the protease, or substituted for one or more amino acid residues within, or at either or both ends of the amino acid sequence of the parent protease.

10

15

20

25

35

The term "isolated nucleic acid sequence" as used herein refers to a nucleic acid sequence which is essentially free of other nucleic acid sequences, e.g., at least about 20% pure, preferably at least about 40% pure, more preferably at least about 60% pure, even more preferably at least about 80% pure, and most preferably at least about 90% pure determined by agarose electrophoresis. For example, isolated nucleic acid sequence can be obtained by standard cloning procedures used in genetic engineering to relocate the nucleic acid sequence from its natural location to a different site where it will be reproduced. The cloning procedures may involve excision and isolation of a desired nucleic acid fragment comprising the nucleic acid sequence encoding the polypeptide, insertion of the fragment into a vector molecule, and incorporation of the recombinant vector into a host cell where multiple copies or clones of the nucleic acid sequence . The nucleic acid sequence may be of will be replicated. genomic, cDNA, RNA, semi synthetic, synthetic origin, or any combinations thereof.

The term "allelic variant" denotes any of two or more alternative forms of a gene occupying the same chromosomal locus. Allelic variation arises naturally through mutation, and may result in polymorphism within populations. Gene mutations can be silent (no change in the encoded polypeptide) or may encode polypeptides having altered amino

عند المناه

acid sequences. The allelic variant of a polypeptide is a polypeptide encoded by an allelic variant of a gene.

The term "nucleic acid construct" is defined herein as a nucleic acid molecule, either single- or double-stranded, which is isolated from a naturally occurring gene or which has been modified to contain segments of nucleic acid which are combined and juxtaposed in a manner which would not otherwise exist in nature. The term nucleic acid construct is synonymous with the term expression cassette when the nucleic acid construct contains all the control sequences required for expression of a coding sequence of the present invention.

The term "coding sequence" is defined herein as a portion of a nucleic acid sequence, which directly specifies the amino acid sequence of its protein product. The boundaries of the coding sequence are generally determined by a ribosome binding site (prokaryotes) or by the ATG start codon (eukaryotes) located just upstream of the open reading frame at the 5' end of the mRNA and a transcription terminator sequence located just downstream of the open reading frame at the 3' end of the mRNA. A coding sequence can include, but is not limited to, DNA, cDNA, and recombinant nucleic acid sequences.

25

10

NOMENCLATURE OF AMINO ACIDS

A	=	Ala	=	Alanine		
V	=	Val	=	Valine		
L	=	Leu	=	Leucine		
I	=	Ile	=	Isoleucine		
P	=	Pro	=	Proline		
F	=	Phe	=	Phenylalanine		
W	=	Trp	=	Tryptophan		
M	=	Met	=	Methionine		
G	=	Gly	=	Glycine		
S	=	Ser	=	Serine		
T	=	Thr	=	Threonine		
С		Cys	=	Cysteine		
Y	=	Tyr = Tyrosine		Tyrosine		
N	=	Asn	=	Asparagine		
Q	=	Gln	=	Glutamine		
D	=	Asp	=	Aspartic Acid		
E	=	Glu	=	Glutamic Acid		
K	=	Lys	=	Lysine		
R	=	Arg	=	Arginine		
Н	=	His	=	Histidine		
X	=	Xaa	=	Any amino acid		
	V L I P F W M G S T C Y N Q D E K R H	V = L = I = I = I = I = I = I = I = I = I	V = Val L = Leu I = Ile P = Pro F = Phe W = Trp M = Met G = Gly S = Ser T = Thr C = Cys Y = Tyr N = Asn Q = Gln D = Asp E = Glu K = Lys R = Arg H = His	V = Val = Leu = I = Ile = P = Pro = F = Phe = W = Trp = M = Met = G = Gly = S = Ser = T = Thr = C = Cys = Y = Tyr = N = Asn = Q = Gln = D = Asp = E = Glu = K = Lys = H = His =		

NOMENCLATURE OF NUCLEIC ACIDS

A	=	Adenine							
G	=	Guanine							
С	=	Cytosine							
T	=	Thymine (only in DNA							
U	=	Uracil	(only	in	RNA)				
N	=	A, C, G or T; A or G;							
R	=								
Y	=	C or T;							
D	=	A, G or T;							
X	=	deoxyinosine.							
	G C T U N R Y	G = C = T = U = N = R = Y = D =	G = Guanine C = Cytosine T = Thymine U = Uracil N = A, C, G or C R = A or G; Y = C or T; D = A, G or T;	G = Guanine C = Cytosine T = Thymine (only U = Uracil (only N = A, C, G or T; R = A or G; Y = C or T; D = A, G or T;	G = Guanine C = Cytosine T = Thymine (only in U = Uracil (only in N = A, C, G or T; R = A or G; Y = C or T; D = A, G or T;				

.. NAMING OF RP-II PROTEASES

In describing the RP-II proteases of the invention the following abbreviations are used for ease of reference:

BLC = RP-II protease from Bacillus licheniformis (cf. US

40 Patent No. 4,266,031),

AA513 = RP-II protease from Bacillus halmapalus AA513

AC116 = RP-II protease from Bacillus licheniformis AC116

BO32 = RP-II protease from Bacillus pumilus BO32

CDJ31 = RP-II protease from Bacillus licheniformis CDJ31

JA96 = RP-II protease from Bacillus pumilus JA96

MPR = RP-II protease from Bacillus subtilis IS75 (cf. EP 369
817 B1)

SEQUENCE LISTING

25

30

35

40

In the appended Sequence Listing the RP-II proteases are indicated as:

SEQ. ID. NO. 1 = BLC (DNA), SEQ. ID. NO. 2 = BLC (AA),

SEQ. ID. NO. 3 = AA513 (DNA), SEQ. ID. NO. 4 = AA513 (AA),

SEQ. ID. NO. 5 = AC116 (DNA), SEQ. ID. NO. 6 = AC116 (AA)

15 SEQ. ID. NO. 7 = BO32 (DNA), SEQ. ID. NO. 8 = BO32 (AA)

SEQ. ID. NO. 9 = CDJ31 (DNA), SEQ. ID. NO. 10 = CDJ31 (AA)

SEQ. ID. NO. 11 = JA96 (DNA), SEQ. ID. NO. 12 = JA96 (AA)

SEQ. ID. NO. 13 = BSMPR (DNA), SEQ. ID. NO. 14 = BSMPR (AA)

20 NOMENCIATURE AND CONVENTIONS FOR DESIGNATION OF VARIANTS

In describing the various enzyme variants produced or contemplated according to the invention, the following nomenclatures and conventions have been adapted for ease of reference:

Aligning the amino acid sequence of a novel isolated or parent wild type enzyme with a suitable well-known enzyme of the same group or class of enzymes first defines a frame of reference. If nothing else is indicated herein, in the present instance the *Bacillus licheniformis* RP-II protease, first designated component C and therefore here abbreviated BLC, has been chosen as standard.

The alignment can be obtained by the GAP routine of the GCG package version 9.1 to number the variants using the following parameters: gap creation penalty = 8 and gap extension penalty = 8 and all other parameters kept at their default values.

This will define a number of deletions and insertions in relation to the standard, here BLC. In the alignments deletions are indicated by asterixes (*) in the referenced sequence, and the referenced enzyme will be considered to

have a gap at the position in question. Insertions are indicated by asterixes (*) in the sequence of the standard enzyme, and the positions in the referenced enzyme are given as the position number of the last amino acid residue where a corresponding amino acid residue exists in the standard enzyme with a lower case letter appended in alphabetical order, e.g. 82a, 82b, 82c, 82d, see Fig. 1

In case the referenced enzyme contains a N- or C-terminal extension in comparison to the standard enzyme, an N-terminal extension is given the position number 0a, 0b, etc.

A C-terminal extension will be given either the position number of the C-terminal amino acid residue of the standard enzyme with a lower case letter appended in alphabetical order, or simply a continued consecutive numbering.

The various modifications performed in a wild type enzyme is indicated in general using three elements as follows:

20

25

30

10

15

Original amino acid Position Substituted amino acid

The notation E152G thus means a substitution of a glutamic acid in position 152 with a glycine.

In the case when the original amino acid residue may be any amino acid residue, a short hand notation may at times be used indicating only the position and substituted amino acid,

Position Substituted amino acid

Such a notation is particular relevant in connection with modification(s) in homologous RP-II proteases.

Similarly when the identity of the substituting amino acid residue(s) is immaterial,

35 Original amino acid Position

When both the original amino acid(s) and substituted amino acid(s) may comprise any amino acid, then only the position is indicated, e.g.: 152.

When the original amino acid(s) and/or substituted
40 amino acid(s) may comprise more than one, but not all amino

acid(s), then the selected amino acids are indicated inside
brackets {},

For specific variants the specific three or one letter codes are used, including the codes Xaa and X to indicate any amino acid residue.

10 SUBSTITUTIONS:

The substitution of Alanine for Glutamic acid in position 152 is designated as:

Glu152Ala or E152A

or the substitution of any amino acid residue acid for Glutamic acid in position 152 is designated as:

Glu152Xaa or E152X

or

15

Glu152 or E152

The substitution of Glutamic acid for any amino acid residue in position 89 would thus be designated

Xaa89Glu or X89E.

or

30

35

89Gluor 89E

Such a notation is particular relevant in connection with modification(s) in homologous RP-II proteases (vide infra). 89Glu is thus meant to comprise e.g. both an Arg89Glu modification in BLC and an Asn89Glu modification in JA96 (cf. Fig. 1b).

For a modification where the original amino acid(s) and/or substituted amino acid(s) may comprise more than one, but not all amino acid(s), the substitution of glycine, alanine, serine or threonine for arginine in position 152 would be indicated by

Glu152{Gly,Ala,Ser,Thr} or E152{G,A,S,T} to indicate the variants

E152G, E152A, E152S, and E152T.

DELETIONS:

A deletion of Glutamic acid in position 152 will be indicated by:

Glu152* or E152*

Correspondingly the deletion of more than one amino acid residue, such as the deletion of glycine and leucine in positions 152 and 153 will be designated

Glu152*+Thr153* or E152*+T153*

INSERTIONS:

15

The insertion of an additional amino acid residue such as e.g. a lysine after E152 is:

Glu152GluLys or E152EK; or when more than one amino acid residue is inserted, such as e.g. a Lys, Ala and Ser after E152 this is:

Glu152GluLysAlaSer or E152EKAS (SEQ ID NO:15)

In such cases the inserted amino acid residue(s) are numbered by the addition of lower case letters to the position number of the amino acid residue preceding the inserted amino acid residue(s). In the above example the sequences 151 to 153 would thus be:

151 152 153

Parent S - E - T

25 151 152 152a 152b 152c 153

Variant S - E - K - A - S - T (SEQ ID NO:16)

In cases where an amino acid residue identical to the existing amino acid residue is inserted it is clear that degeneracy in the nomenclature arises. If for example a glutamic acid is inserted after the glutamic acid in the above example this would be indicated by E152EE. The same actual change could just as well be indicated as S151SE for the change from

35 151 152 153

Parent S - E - T

to

14

151 152 152a 153

Variant S - E - E - T (SEQ ID NO:17)

151 151a 152 153

Such instances will be apparent to the skilled person, and the indication E152EE and corresponding indications for this type of insertions are thus meant to comprise such equivalent degenerate indications.

Correspondingly the modification of a residue and
simultaneous insertion of a further residue may be designated
in different ways as

V110PS = V110VS+V110P = V110P+P110PS indicating that the position 110 valine has been substituted by a proline and a serine.

15

20

FILLING A GAP:

Where a deletion in an enzyme exists in the reference comparison with the standard sequence used for the numbering, an insertion in such a position is indicated as:

*121Ser or *121S for the insertion of an serine in position 121

MULTIPLE MODIFICATIONS

Variants comprising multiple modifications are separated by pluses, e.g.:

Ser1Val+Glu152Ala or S1V+E152A representing modifications in positions 1 and 152 substituting serine and glutamic acid for valine and alanine, respectively.

Or e.g. Arg8{Gly,Ala,Ser,Thr}+Glu152{Gly,Ala,Ser,Thr} designates the variants

Arg8Ala+Glu152Gly, Arg8Gly+Glu152Gly, Arg8Ser+Glu152Gly, Arg8Thr+Glu152Gly, Arg8Gly+Glu152Ala, . Arg8Ala+Glu152Ala, Arg8Thr+Glu152Ala, 35 Arg8Ser+Glu152Ala, Arg8Ala+Glu152Ser, __ Arg8Gly+Glu152Ser, Arg8Ser+Glu152Ser, Arg8Thr+Glu152Ser, Arg8Gly+Glu152Thr, Arg8Ala+Glu152Thr, Arg8Ser+Glu152Thr, and Arg8Thr+Glu152Thr.

This nomenclature is particularly relevant in relation to modifications aimed at substituting, replacing, inserting or deleting amino acid residues having specific common properties, such as residues of positive charge (K, R, H), negative charge (D, E), or conservative amino acid modification(s) of e.g.

Arg8{Glu,Asp,Lys}+Glu152{Asp,Arg,Lys}, which signifies substituting a charged amino acid for another charged amino acid. See section "Detailed description of the invention" for further details.

Proteases

10

20

25

35

40

Enzymes cleaving the amide linkages in protein substrates are classified as proteases, or (interchangeably) peptidases (see Walsh, 1979, Enzymatic Reaction Mechanisms. W.H. Freeman and Company, San Francisco, Chapter 3).

Serine proteases

A serine protease is an enzyme which catalyzes the hydrolysis of peptide bonds, and in which there is an essential serine residue at the active site (White, Handler and Smith, 1973 "Principles of Biochemistry," Fifth Edition, McGraw-Hill Book Company, NY, pp. 271-272).

The bacterial serine proteases have molecular weights in the 20,000 to 45,000 Dalton range. They are inhibited by disopropylfluorophosphate. They hydrolyze simple terminal esters and are similar in activity to eukaryotic chymotrypsin, also a serine protease.

30 Description of the RP-II protease from B. licheniformis ATCC 14580

For ease of reference, the following disclosure of recombinantly produced RP-II variants is based on the RP-II protease derived from the species B. licheniformis ATCC 14580, the amino acid sequence of which is shown in Fig. 1 below. It will be understood, however, that also functional analogues of RP-II proteases derivable from other Bacilli, such as the novel wild type RP-II proteases disclosed herein, may be modified in a manner similar to that described herein for the B. licheniformis ATCC 14580—RP-II protease. Accordingly,

variants of such functional analogous are considered to be within the scope of the present invention. Examples of other Bacillus strains, which have been found to produce RP-II proteases, are Bacillus pumilus, Bacillus subtilis, and Bacillus halmapalus. However, it is expected that RP-II proteases will be found in many more Bacillus strains.

The parent B. licheniformis RP-II protease, BLC, as disclosed in US Patent No. 4,266,031 has the amino acid sequence shown in Fig. 1 and SEQ ID NO: 2, and the corresponding DNA sequence is shown in SEQ ID NO: 1.

DETAILED DESCRIPTION OF THE INVENTION

The invention will be disclosed in detail in the following sections.

15

20

25

30

35

40

Novel Isolated Rp-II Proteases And Nucleic Acid Sequences Encoding These

Accordingly, in a first embodiment the present invention relates to novel isolated RP-II proteases that are immunochemically identical or partially identical by cross-reaction with an antibody raised against or reactive with at least one epitope of a RP-II protease comprising the amino acid sequences of the mature peptides shown in the appended Sequence Listing ID No. 4, ID No. 6, ID No. 8, ID No. 10, or ID No. 12.

The immunological cross reactivity, may be assayed using an antibody raised against or reactive with at least one epitope of a RP-II protease comprising the amino acid sequence of the mature peptide shown in SEQ ID NO: 2, 4, 6, 8, 10, and antibody, which may either be monoclonal polyclonal, may be produced by methods known in the art, e.g. as described by Hudson et al., 1989, Practical Immunology, 3. Blackwell Scientific Publications. The immunological cross-reactivity may be determined using assays known in the art, examples of which are Western Blotting or radial immunodiffusion assay, e.g. as described by Hudson et al., 1989. According to such assays the polypeptides of the invention can be characterized as being partially " immunochemically identical, or preferably immunochemically

_ <u>:----------</u>

identical to each other. The immunochemical properties can furthermore be determined immunologically by cross-reaction identity tests. The identity tests can be performed by the well-known Ouchterlony double immunodiffusion procedure or by tandem crossed immunoelectrophoresis according to N. H. Axelsen; Handbook of Immunoprecipitation-in-Gel Techniques; Blackwell Scientific Publications (1983), chapters 5 and 14. The terms "antigenic identity" and "partial antigenic identity" are described in the same book, chapters 5, 19 and 20.

10

15

20

25

30

35

40

In a second embodiment, the present invention relates to isolated RP-II proteases having an amino acid sequence which has a degree of identity to the mature polypeptides of SEQ ID NOs: 2, 4, 6, 8, 10 or 12 of at least about 60%, preferably at least about 70%, more preferably at least about 80%, even more preferably at least about 90%, most preferably at least about 95%, and even most preferably at least about 97%, which have protease activity (hereinafter "homologous RP-II proteases"). In a preferred embodiment, the homologous RP-II proteases have an amino acid sequence which differs by five amino acids, preferably by four amino acids, more preferably by three amino acids, even more preferably by two amino acids, and most preferably by one amino acid from the amino acid sequence of the mature polypeptides of SEQ ID NO: 2, 4, 6, 8, 10 or 12. For purposes of the present invention, the degree of identity between two amino acid sequences is determined by the GAP method described above.

Preferably, RP-II proteases of the present invention comprise the amino acid sequence of SEQ ID NOs: 2, 4, 6, 8, 10, or 12 or allelic variants thereof; or a fragment thereof that has protease activity. In a more preferred embodiment, the RP-II proteases of the present invention comprise the amino acid sequence of SEQ ID NOs: 2, 4, 6, 8, 10 or 12. In another preferred embodiment, the RP-II proteases of the present invention comprise the amino acid sequences of the mature polypeptides of SEQ ID NO: 2, 4, 6, 8, 10, or 12, or allelic variants thereof; or a fragment thereof that has protease activity. In another preferred embodiment, the RP-II proteases of the present invention comprise the amino acid sequences of the mature polypeptides of SEQ=ID NO: 2, 4, 6,

In another preferred embodiment, the RP-II 8, 10, or 12. proteases of the present invention consist of the amino acid sequences of SEQ ID NO: 2, 4, 6, 8, 10, or 12 or an allelic variant thereof; or a fragment thereof, wherein the fragment has protease activity. In another preferred embodiment, the RP-II proteases of the present invention consist of the amino acid sequence of SEQ ID NOs: 2, 4, 6, 8, 10, or 12. another preferred embodiment, RP-II proteases of the present invention consists of the amino acid sequences of the mature polypeptides of SEQ ID NO: 2, 4, 6, 8, 10, or 12, or an allelic variant thereof; or a fragment thereof that has protease activity. In another preferred embodiment, the RP-II proteases of the present invention consist of the amino acid sequences of the mature polypeptides of SEQ ID NO: 2, 4, 6, 8, 10, or 12.

10

15

20

25

30

35

In a third embodiment, the present invention relates to isolated RP-II proteases encoded by nucleic acid sequences which hybridize under very low stringency conditions, preferably low stringency conditions, more preferably medium stringency conditions, more preferably medium-high stringency conditions, even more preferably high stringency conditions, and most preferably very high stringency conditions with a nucleic acid probe which hybridizes under the same conditions with (i) a nucleic acid sequence encoding a RP-II protease of SEQ ID NO: 2, 4, 6, 8, 10, 12, or 14, (ii) the cDNA sequence encoding a RP-II protease of SEQ ID NO:2, 4, 6, 8, 10, 12, or (iii) a subsequence of (i) or (ii), or complementary strand of (i), (ii), or (iii) (J. Sambrook, E.F. Fritsch, and T. Maniatus, 1989, Molecular Cloning, A Laboratory Manual, 2d edition, Cold Spring Harbor, New York). The subsequence of SEQ ID NO: 2, 4, 6, 8, 10, 12, or 14 may be at least 100 nucleotides or preferably at least 200 nucleotides. Moreover, the subsequence may encode a RP-II protease fragment that has protease activity.

In a fourth embodiment, the present invention relates to isolated RP-II proteases encoded by nucleic acid sequences which hybridize under very low stringency conditions, preferably low stringency conditions, more preferably medium stringency conditions, more preferably medium-high stringency conditions, even more preferably—high stringency conditions,

and most preferably very high stringency conditions with a

nucleic acid probe which hybridizes under the same conditions

with (i) the nucleic acid sequence of SEQ ID NO:1, 3, 5, 7, 9, 11, or 13, (ii) the cDNA sequence of SEQ ID NO:1, 3, 5, 7,

9, 11, 13 (iii) a subsequence of (i) or (ii), or (iv) a

complementary strand of (i), (ii), or (iii) (J. Sambrook,

E.F. Fritsch, and T. Maniatus, 1989, Molecular Cloning, A

Laboratory Manual, 2d edition, Cold Spring Harbor, New York).
The subsequence of SEQ ID NO: 1, 3, 5, 7, 9, 11, or 13 may

be at least 100 nucleotides or preferably at least 200

nucleotides. Moreover, the subsequence may encode a

polypeptide fragment that has protease activity.

10

15

20

25

30

35

40

The nucleic acid sequence of SEQ ID NO: 1, 3, 5, 7, 9, 11, or 13 or a subsequence thereof, as well as the amino acid sequence of SEQ ID NO: 2, 4, 6, 8, 10 12, or 14 or a fragment thereof, may be used to design a nucleic acid probe to identify and clone DNA encoding RP-II proteases from strains of different genera or species according to methods well known in the art. In particular, such probes can be used for hybridization with the genomic or cDNA of the genus or species of interest, following standard Southern blotting order to identify and isolate procedures, in corresponding gene therein. Such probes can be considerably shorter than the entire sequence, but should be at least 15, preferably at least 25, and more preferably at least 35 nucleotides in length. Longer probes can also be used. Both DNA and RNA probes can be used. The probes are typically labelled for detecting the corresponding gene (for example, with 32P, 3H, 35S, biotin, or avidin). Such probes are encompassed by the present invention.

Thus, a genomic DNA or cDNA library prepared from such other organisms may be screened for DNA that hybridizes with the probes described above and which encodes a RP-II protease. Genomic or other DNA from such other organisms may be separated by agarose or polyacrylamide gel electrophoresis, or other separation techniques. DNA from the libraries or the separated DNA may be transferred to and immobilized on nitrocellulose or other suitable carrier material. In order to identify a clone or DNA which is homologous with SEQ ID NO: 1, 3, 5, 7, 9, 11, or 13, or a

<u>_</u>_____

subsequence thereof, the carrier material is used in a Southern blot. For purposes of the present invention, hybridization indicates that the nucleic acid sequence hybridizes to a nucleic acid probe corresponding to the nucleic acid sequence shown in SEQ ID NO: 1, 3, 5, 7, 9, 11, or 13, its complementary strand, or a subsequence thereof, under very low to very high stringency conditions. Molecules to which the nucleic acid probe hybridizes under these conditions are detected using X-ray film.

For long probes of at least 100 nucleotides in length, very low to very high stringency conditions are defined as prehybridization and hybridization at 42°C in 6XSSC, 5x Denhardt's solution, 0.2% SDS, 100 mg/ml sheared and denatured salmon sperm DNA, and either 25% formamide for very low and low stringencies, 35% formamide for medium and medium-high stringencies, or 50% formamide for high and very high stringencies, following standard Southern blotting procedures.

10

15

20

25

30

35

For long probes of at least 100 nucleotides in length, the carrier material is finally washed three times each for 15 minutes using 2 x SSC, 0.2% SDS preferably at least at 45°C (very low stringency), more preferably at least at 50°C (low stringency), more preferably at least at 55°C (medium stringency), more preferably at least at 60°C (medium-high stringency), even more preferably at least at 65°C (high stringency), and most preferably at least at 70°C (very high stringency).

For short probes or synthetic oligonucleotides probes which are about 15 nucleotides to about 30 nucleotides in length, stringency conditions are defined as prehybridization, hybridization, and washing posthybridization at 5°C to 10°C below the calculated Tm using the calculation according to Bolton and McCarthy (1962, Proceedings of the National Academy of Sciences USA 48:1390) 5 x Denhardt's solution, 0.05% x SSC. pyrophosphate, 100 mg/ml sheared and denatured herring sperm SDS following standard Southern blotting DNA. 0.5% procedures.

For short end=labelled 32P probes or synthetic _ 40 oligonucleotides end-labelled 32P probes which are about 15

nucleotides to about 30 nucleotides in length, the carrier material is washed in prewarmed 6X SCC plus 0.05% sodium pyrophosphate for 15 to 30 minutes at 5°C to 10°C below the calculated Tm. The wash is repeated until a Geiger counter is not exhibiting above background radioactivity.

In a further embodiment, the present invention relates to isolated nucleic acid sequences encoding RP-II proteases having an amino acid sequence which has a degree of identity to the mature peptides of SEQ ID NOs: 2, 4, 6, 8, 10 or 12 of at least about 60%, preferably at least about 70%, more preferably at least about 80%, even more preferably at least about 90%, most preferably at least about 95%, and even most preferably at least about 97%, which have protease activity (hereinafter "homologous RP-II proteases"). In a preferred embodiment, the homologous RP-II proteases have an amino acid sequence which differs by five amino acids, preferably by four amino acids, more preferably by three amino acids, even more preferably by two amino acids, and most preferably by one amino acid from the amino acid sequence of the mature polypeptides of SEQ ID NO: 2, 4, 6, 8, 10 or 12. purposes of the present invention, the degree of identity between two amino acid sequences is determined by the GAP method described above.

10

15

20

25

30

35

40

Preferably, the nucleic acid sequences of the present invention encode RP-II proteases that comprise the amino acid sequence of SEQ ID NOs: 2, 4, 6, 8, 10, or 12 or allelic variants thereof; or a fragment thereof that has protease activity. In a more preferred embodiment, the nucleic acid sequence of the present invention encodes a RP-II protease that comprises the amino acid sequence of SEQ ID NOs: 2, 4, 6, 8, 10 or 12. In another preferred embodiment, the nucleic acid sequence of the present invention encodes a RP-II protease that comprises an amino acid sequence of the mature polypeptides of SEQ ID NO: 2, 4, 6, 8, 10, or 12, or allelic variants thereof; or a fragment thereof that has protease In another preferred embodiment, the nucleic acid sequence of the present invention encodes a RP-II protease that _comprises the amino acid sequence of a mature polypeptide of SEQ ID NO: 2, 4, 6, 8, 10, or 12._ In another preferred embodiment, the nucleic acid sequence of the

present invention encodes a RP-II protease that consists of the amino acid sequence of SEQ ID NO: 2, 4, 6, 8, 10, or 12, or an allelic variant thereof; or a fragment thereof, wherein In another the polypeptide fragment has protease activity. preferred embodiment, the nucleic acid sequence of the present invention encodes a RP-II protease that consists of the amino acid sequence of SEQ ID NO: 2, 4, 6, 8, 10, or 12. In another preferred embodiment, the nucleic acid sequence of the present invention encodes a RP-II protease that consists of the amino acid sequence of a mature polypeptide of SEQ ID NO: 2, 4, 6, 8, 10, or 12, or an allelic variant thereof; or a fragment thereof that has protease activity. In another preferred embodiment, the nucleic acid sequence of the present invention encodes a RP-II protease that consists of amino acid sequence of a mature polypeptide of SEQ ID NO: 2, 4, 6, 8, 10, or 12.

10

15

20

25

30

35

40

The present invention also encompasses nucleic acid sequences which encode a RP-II protease having the amino acid sequence of SEQ ID NO: 2, 4, 6, 8, 10, or 12, which differ from SEQ ID NO: by virtue of the degeneracy of the genetic code. The present invention also relates to subsequences of SEQ ID NO: 1, 3, 5, 7, 9, or 11 which encode fragments of SEQ ID NO: 2, 4, 6, 8, 10, or 12 which have protease activity.

A subsequence of SEQ ID NO: 1, 3, 5, 7, 9, or 11 is a nucleic acid sequence encompassed by SEQ ID NO: 1, 3, 5, 7, 9, or 11 except that one or more nucleotides from the 5' and/or 3' end have been deleted.

In a yet further embodiment, the present invention relates to isolated nucleic acid sequences encoding RP-II proteases of the invention which hybridize under very low stringency conditions, preferably low stringency conditions, more preferably medium stringency conditions, more preferably medium-high stringency conditions, even more preferably high and most preferably very conditions, stringency stringency conditions with a nucleic acid probe which hybridizes under the same conditions with (i) the nucleic acid sequence of SEQ ID NO: 1, 3, 5, 7, 9, or 11, (ii) the cDNA sequence of SEQ ID NO:1, (iii) a subsequence of (i) or (ii), or (iv) a complementary strand of (i), (ii), or (iii) (J. Sambrook, E.F. Fritsch, and T. Maniatus, 1989, Molecular

Cloning, A Laboratory Manual, 2d edition, Cold Spring Harbor, New York). The subsequence of SEQ ID NO: 1, 3, 5, 7, 9, or 11 may be at least 100 nucleotides or preferably at least 200 nucleotides. Moreover, the subsequence may encode a polypeptide fragment that has protease activity.

The polypeptides encoded by the isolated nucleic acid sequences of the present invention have at least 20%, preferably at least 40%, more preferably at least 60%, even more preferably at least 80%, even more preferably at least 90%, and most preferably at least 100% of the protease activity of the mature RP-II proteases of SEQ ID NO: 2, 4, 6, 8, 10, or 12.

15 Description of RP-II protease variants of the invention

In the following specific classes of RP-II protease variants of the invention having improved properties are described as well as the concepts used for the design of such variants.

20

25

30

35

10

Stabilization by modification of Asn-Gly Pairs

It is known that at alkaline pH, the side chain of Asn may interact with the NH group of a sequential neighbouring amino acid to form an isoAsp residue where the backbone goes through the Asp side chain. This will leave the backbone more vulnerable to proteolysis. The deamidation is much more likely to occur if the residue that follows is a Gly. Changing the Asn in front of the Gly or the Gly will prevent this from happening and thus improve the stability, especially as concerns thermo- and storage stability.

The invention consequently further relates to a RP-II protease variant, in which either or both residues of any of the Asn-Gly sequence appearing in the amino acid sequence of the parent RP-II protease is/are deleted or substituted with a residue of a different amino acid.

The Asn and/or Gly residue may, for instance, be substituted with a residue of an amino acid selected from the group consisting of A, Q, S, P, T and Y.

of the BLC protease; positions 68-69, and/or 192-193 of the AC116 protease, positions 68-69 and/or 192-193 of the CDJ-31 protease, positions 45-46, 74-75, 187-188, and/or 191-192 of the BO32 protease, positions 45-46, 74-75, 187-188 and/or 191-192 of the JA96 protease, and positions 90-91, and/or 195-196 of the AA513 protease, and positions 78-79, 103-104 and or 192a-192b of the MPR protease may be deleted or substituted with a residue of an amino acid selected from the group consisting of A, Q, S, P, T and Y.

10

```
Specific variants of BLC are:
```

 $N68\{*,A,Q,S,P,T,Y\};$ $G69\{*,A,Q,S,P,T,Y\}$

 $N68\{*,A,Q,S,P,T,Y\}+G69\{*,A,Q,S,P,T,Y\}$

N182{*,A,Q,S,P,T,Y}; G183{*,A,Q,S,P,T,Y}

15 N182{*,A,Q,S,P,T,Y}+G183{*,A,Q,S,P,T,Y}

 $N192\{*,A,Q,S,P,T,Y\};$ $G193\{*,A,Q,S,P,T,Y\}$

 $N192\{*,A,Q,S,P,T,Y\}+G193\{*,A,Q,S,P,T,Y\}$

 $N68\{*,A,Q,S,P,T,Y\}+N182\{*,A,Q,S,P,T,Y\}$

 $N68\{*,A,Q,S,P,T,Y\}+N182\{*,A,Q,S,P,T,Y\}+N192\{*,A,Q,S,P,T,Y\}$

20

Specific variants of the AC116 protease are:

 $N68\{*,A,Q,S,P,T,Y\};$ $G69\{*,A,Q,S,P,T,Y\}$

 $N68\{*,A,Q,S,P,T,Y\}+G69\{*,A,Q,S,P,T,Y\}$

 $N192\{*,A,Q,S,P,T,Y\};$ $G193\{*,A,Q,S,P,T,Y\}$

25 $N192\{*,A,Q,S,P,T,Y\}+G193\{*,A,Q,S,P,T,Y\}$

 $N68\{*,A,Q,S,P,T,Y\}+N192\{*,A,Q,S,P,T,Y\}$

Specific variants of CDJ-31 are:

 $N68\{*,A,Q,S,P,T,Y\};$ $G69\{*,A,Q,S,P,T,Y\}$

30 N68 $\{*,A,Q,S,P,T,Y\}+G69\{*,A,Q,S,P,T,Y\}$

 $N192\{*,A,Q,S,P,T,Y\};$ $G193\{*,A,Q,S,P,T,Y\}$

 $N192\{*,A,Q,S,P,T,Y\}+G193\{*,A,Q,S,P,T,Y\}$

 $N68\{*,A,Q,S,P,T,Y\}+N192\{*,A,Q,S,P,T,Y\}$

35 Specific variants of BO32 are:

 $N45\{*,A,Q,S,P,T,Y\};$ $G46\{*,A,Q,S,P,T,Y\}$

 $N45\{*,A,Q,S,P,T,Y\}+G46\{*,A,Q,S,P,T,Y\}$

 $N74\{*,A,Q,S,P,T,Y\};$ $G75\{*,A,Q,S,P,T,Y\}$

 $N74\{*,A,Q,S,P,T,Y\}+G75\{*,A,Q,S,P,T,Y\}$

40 N187{*,A,Q,S,P,T,Y}; G188{*,A,Q,S,P,T,Y}

```
N187\{*,A,Q,S,P,T,Y\}+G188\{*,A,Q,S,P,T,Y\}
    N192{*,A,Q,S,P,T,Y};
                             G193{*,A,Q,S,P,T,Y}
    N192\{*,A,Q,S,P,T,Y\} + G193\{*,A,Q,S,P,T,Y\}
    N45\{*,A,Q,S,P,T,Y\}+N74\{*,A,Q,S,P,T,Y\}
 5 N45{*,A,Q,S,P,T,Y}+N187{*,A,Q,S,P,T,Y}
    N45\{*,A,Q,S,P,T,Y\}+N192\{*,A,Q,S,P,T,Y\}
    N74\{*,A,Q,S,P,T,Y\}+N187\{*,A,Q,S,P,T,Y\}
    N74\{*,A,Q,S,P,T,Y\}+N192\{*,A,Q,S,P,T,Y\}
    N187\{*,A,Q,S,P,T,Y\}+N192\{*,A,Q,S,P,T,Y\}
    N45{*,A,Q,S,P,T,Y}+N74{*,A,Q,S,P,T,Y}+N187{*,A,Q,S,P,T,Y}
10
    N45\{*,A,Q,S,P,T,Y\}+N74\{*,A,Q,S,P,T,Y\}+N192\{*,A,Q,S,P,T,Y\}
    N45\{*,A,Q,S,P,T,Y\}+N187\{*,A,Q,S,P,T,Y\}+N192\{*,A,Q,S,P,T,Y\}
    N74\{*,A,Q,S,P,T,Y\}+N187\{*,A,Q,S,P,T,Y\}+N192\{*,A,Q,S,P,T,Y\}
    N45\{*,A,Q,S,P,T,Y\}+N74\{*,A,Q,S,P,T,Y\}+N187\{*,A,Q,S,P,T,Y\}+N192
    {*,A,Q,S,P,T,Y}
15
    Specific variants of JA96 are:
    N45{*,A,Q,S,P,T,Y};
                              G46{*,A,Q,S,P,T,Y}
    N45\{*,A,Q,S,P,T,Y\}+G46\{*,A,Q,S,P,T,Y\}
20 N74{*,A,Q,S,P,T,Y};
                              G75{*,A,Q,S,P,T,Y}
    N74\{*,A,Q,S,P,T,Y\}+G75\{*,A,Q,S,P,T,Y\}
    N187{*,A,Q,S,P,T,Y};
                             G188{*,A,Q,S,P,T,Y}
    N187{*,A,Q,S,P,T,Y}+G188{*,A,Q,S,P,T,Y}
    N192\{*,A,Q,S,P,T,Y\}; G193\{*,A,Q,S,P,T,Y\}
    N192\{*,A,Q,S,P,T,Y\} + G193\{*,A,Q,S,P,T,Y\}
    N45\{*,A,Q,S,P,T,Y\}+N74\{*,A,Q,S,P,T,Y\}
    N45\{*,A,Q,S,P,T,Y\}+N187\{*,A,Q,S,P,T,Y\}
    N45\{*,A,Q,S,P,T,Y\}+N192\{*,A,Q,S,P,T,Y\}
    N74\{*,A,Q,S,P,T,Y\}+N187\{*,A,Q,S,P,T,Y\}
    N74\{*,A,Q,S,P,T,Y\}+N192\{*,A,Q,S,P,T,Y\}
30
    N187\{*,A,Q,S,P,T,Y\}+N192\{*,A,Q,S,P,T,Y\}
    N45{*,A,Q,S,P,T,Y}+N74{*,A,Q,S,P,T,Y}+N187{*,A,Q,S,P,T,Y}
    N45\{*,A,Q,S,P,T,Y\}+N74\{*,A,Q,S,P,T,Y\}+N192\{*,A,Q,S,P,T,Y\}
    N45\{*,A,Q,S,P,T,Y\}+N187\{*,A,Q,S,P,T,Y\}+N192\{*,A,Q,S,P,T,Y\}
    N74\{*,A,Q,S,P,T,Y\}+N187\{*,A,Q,S,P,T,Y\}+N192\{*,A,Q,S,P,T,Y\}
    N45\{*,A,Q,S,P,T,Y\}+N74\{*,A,Q,S,P,T,Y\}+N187\{*,A,Q,S,P,T,Y\}+N192
     {*,A,Q,S,P,T,Y}
```

Specific variants of AA513 are:

40 N90 $\{*,A,Q,S,P,T,Y\}$; = G91 $\{*,A,Q,S,P,T,Y\}$

```
N90{*,A,Q,S,P,T,Y}+G91{*,A,Q,S,P,T,Y}
N195{*,A,Q,S,P,T,Y}; G196{*,A,Q,S,P,T,Y}
N195{*,A,Q,S,P,T,Y}+G196{*,A,Q,S,P,T,Y}
N90{*,A,Q,S,P,T,Y}+N195{*,A,Q,S,P,T,Y}
```

5

20

25

30

35

Specific variants of MPR are:

N78{*,A,Q,S,P,T,Y}; G79{*,A,Q,S,P,T,Y} N78{*,A,Q,S,P,T,Y}+G79{*,A,Q,S,P,T,Y} N103{*,A,Q,S,P,T,Y}; G104{*,A,Q,S,P,T,Y} N103{*,A,Q,S,P,T,Y}+G104{*,A,Q,S,P,T,Y}

0 N103{*,A,Q,S,P,T,Y}+G104{*,A,Q,S,P,T,Y}
N192a{*,A,Q,S,P,T,Y}; G192b{*,A,Q,S,P,T,Y}
N192a{*,A,Q,S,P,T,Y}+G192b{*,A,Q,S,P,T,Y}
N78{*,A,Q,S,P,T,Y}+N103{*,A,Q,S,P,T,Y}

 $N78\{*,A,Q,S,P,T,Y\}+N192a7\{*,A,Q,S,P,T,Y\}$

15 N103{*,A,Q,S,P,T,Y}+N192a{*,A,Q,S,P,T,Y} N78{*,A,Q,S,P,T,Y}+N103{*,A,Q,S,P,T,Y}+N192a{*,A,Q,S,P,T,Y}

Removal of autoproteolysis sites

According to a further aspect of the invention autoproteolysis sites may be removed by changing the amino acids at an autoproteolysis site. Since the RP-II proteases cleaves at Glu and Asp residues it is preferred to modify such residues of a parent RP-II protease having the same or a similar specificity, preferably by substituting with any other amino acid except Glu.

The parent RP-II proteases are mostly specific towards Glu and to a minor extend towards Asp residues. Therefore the modification of the parent (trypsin-like) RP-II protease may preferably be made by changing Glu to another amino acid residue (including Asp). Experiments have indicated that the substitution of Ala for Glu or Asp provides good results.

The following Glu and Asp residue positions are found in the BLC protease: E101, E104, E152, E173, E209, D6, D7, D51, D135, D161, D212.

Specific BLC variants are thus E101A, E104A, E152A, E173A, E209A, D6A, D7A, D51A, D135A, D161A, D212A, and double, triple, quadruple, etc. combinations thereof.

<u>.</u>

In JA96 Glu and Asp are found in positions: E81, E143, 40 E151, E202, D5, D6, D69, D96, D103, D135, D152, D161, D173.

Specific JA96 variants are thus E81A, E143A, E151A, E202A, D5A, D6A, D69A, D96A, D103A, D135A, D152A, D161A, D173A, and double, triple, quadruple, etc. combinations thereof.

Corresponding variants are easily identified in any other RP-II protease.

Alternatively autoproteolysis can be prevented by changing the amino acid residue occupying the $1^{\rm st}$ and/or $2^{\rm nd}$ position following the Glu or Asp residue in question to Pro.

c10 For instance, this may in BLC be done in the positions 174 and/or 175 as follows:

Q174P

5

25

30

S175P

0174P+S175P

or in a similar manner in JA96 in positions 152 and/or 153 as D152P; T153P; or D152P+T153P

Corresponding variants are easily identified in any other RP-II protease.

20 Removal of critical oxidation sites

In order to increase the stability of the RP-II protease it may be advantageous to substitute critical oxidation sites, such as methionines, with other amino acid residues which are not subject to oxidation.

Accordingly, in a further embodiment the present invention relates to a RP-II protease variant, in which one or more amino acid residues susceptible to oxidation, especially methionine residues exposed to the surface of the molecule, is/are deleted or replaced with another amino acid residue less susceptible to oxidation. The amino acid residue less susceptible to oxidation may for instance be selected from the group consisting of A, E, N, Q, I, L, S and K.

Specific such variants comprises at least one of the deletions or substitutions

of the BO32 and JA96 proteases, M159{*,S,A,N,Q,K}; of the BO32 protease.

5 Modification of tryptophan residues

In order to stabilize the protein it may be advantageous to replace or delete tryptophan residues at the surface of the protein, e.g. as described in US 5,118,623. The tryptophan residues may advantageously be substituted for F, T, Q or G. Thus, in a further embodiment the invention relates to a RP-II variant comprising one or more of the following substitutions:

```
BLC:
    W35{F,T,Q,G}
15 W88{F,T,Q,G}
    W142\{F,T,Q,G\}
    W217\{F,T,Q,G\}
    AC116:
    W35{F,T,Q,G}
20 W88{F,T,Q,G}
    W142{F,T,Q,G}
    W217\{F,T,Q,G\}
    CDJ31:
    W142\{F,T,Q,G\}
25 W217{F,T,Q,G}
    BO32:
    W142\{F,T,Q,G\}
    JA96:
    W142{F,T,Q,G}
30 AA513:
    W30\{F,T,Q,G\}
    W72{F,T,Q,G}
    W142\{F,T,Q,G\}
    MPR:
```

10

35 W57{F,T,Q,G} W88{F,T,Q,G} W112{F,T,Q,G} W142{F,T,Q,G} W217{F,T,Q,G}

Variants with improved wash performance

The ability of an enzyme to catalyse the degradation of various naturally occurring substrates present on the objects to be cleaned during e.g. wash is often referred to as its washing ability, washability, detergency, or wash performance. The present invention provides novel RP-II proteases for the use in detergents and novel RP-II protease variants exhibiting an improved wash performance as compared to that of the parent RP-II protease.

10 Examples of specific BLC variants comprises one or more of the following substitutions:

 $E152{A,R,K,G}$

E173A

E209A

20

15 E152G+G164R

In relation to wash performance it has been found that the modification of certain tyrosine residues to phenylalanine provides an improved wash performance. Without being bound by any specific theory, it is believed that titration of these Tyr residues in the alkaline wash liquor has negative effects that are alleviated by replacing the Tyr residues with other residues, especially Phe or Trp, particularly Phe.

In the BLC parent RP-II protease, the following tyrosine residues may be modified:

17, 19, 50, 72, 74, 82, 95, 97, 112, 115, 117, 132, 154, 158, 163, 195, 200

Examples of specific BLC variants comprises one or more of the following substitutions:

30 $Y17\{F,W\}$

Y19{F,W}

Y50{F,W}

Y72{F,W}

Y74{F,W}

35 Y82{F,W}

Y95{F,W}

Y97{F,W}

Y112{F,W}

Ý115{F,W}

40 Y117{F,W}

 $Y132\{F,W\}$

Y154{F,W}

Y158{F,W}

Y163{F,W}

5 Y195{F,W}

Y200{F,W}

In the AC116 parent RP-II protease, the following tyrosine residues may be modified:

10 19, 50, 72, 74, 82, 95, 97, 112, 115, 117, 132, 154, 163, 172, 195, 200

Examples of specific AC116 variants comprises one or more of the following substitutions:

Y19{F,W}

15 Y50{F,W}

Y72{F,W}

Y74 {F, W}

Y82{F,W}

Y95{F,W}

20 Y97{F,W}

Y112{F,W}

Y115{F,W}

Y117{F,W}

Y132{F,W}

25 Y154{F,W}

Y158{F,W}

Y163{F,W}

Y172{F,W}

Y195{F,W}

30 Y200{F,W}

In the CDJ31 parent RP-II protease, the following tyrosine residues may be modified:

17, 19, 50, 72, 74, 82, 88, 95, 97, 112, 115, 117, 132, 154,

35 158, 163, 172, 195, 200

__ Examples of specific CDJ31 variants comprises one or more of the following substitutions:

Y17{F,W}__

__Y19{F,W}

-40 Y50{F,W}

```
Y72{F,W}
    Y74{F,W}
    Y82{F,W}
    Y88{F,W}
 5 Y95{F,W}
    Y97{F,W}
    Y112{F,W}
    Y115{F,W}
    Y117{F,W}
10 Y132{F,W}
    Y154{F,W}
    Y158{F,W}
    Y163{F,W}
    Y172{F,W}
15 Y195{F,W}
    Y200{F,W}
```

In the BO32 parent RP-II protease, the following tyrosine residues may be modified:

20 19, 50, 57, 64, 83, 88, 95, 112, 132, 157, 158, 185, 206

Examples of specific BO32 variants comprises one or more of the following substitutions:

Y19{F,W}

Y50{F,W}

25 Y57{F,W}

Y64{F,W}

Y83{F,W}

Y88{F,W}

Y95{F,W}

30 Y112{F,W}

Y132{F,W}

Y157{F,W}

Y158{F,W}

Y185{F,W}

35 Y206{F,W}

In the JA96 parent RP-II protease, the following tyrosine residues may be modified:

_ 19, 24, 50, 57, 64, 83, 88, 95, 112, 132, 157, 158, 186, 206

Examples of specific JA96 variants comprises one or more of the following substitutions:

Y19{F,W}

Y24 {F, W}

5 Y50{F,W}

Y57{F,W}

Y64 {F, W}

Y83{F,W}

Y88{F,W}

10 Y95{F,W}

Y112{F,W}

Y132{F,W}

Y157{F,W}

Y158{F,W}

15 Y186{F,W}

Y206{F,W}

In the AA513 parent RP-II protease, the following tyrosine residues may be modified:

20 24, 74, 77, 84, 88, 97, 130, 132, 167, 172, 186

Examples of specific AA513 variants comprises one or more of the following substitutions:

Y24 {F, W}

Y74 {F, W}

25 Y77{F,W}

Y84 {F, W}

Y88{F,W}

Y87{F,W}

Y97{F,W}

30 Y130{F,W}

Y132{F,W}

 $Y167\{F,W\}$

Y172{F,W}

Y186{F,W}

35

In the MPR parent RP-II protease, the following tyrosine residues may be modified:

19, 26d, 30, 50, 72, 74, 77, 82, 95, 97, 113, 115, 154, 158, 163, 172, 175, 200, 216

Examples of specific MPR variants comprises one or more of the following substitutions:

Y19{F,W}
Y26d{F,W}
5 Y30{F,W}
Y50{F,W}
Y72{F,W}
Y74{F,W}
Y77{F,W}
10 Y82{F,W}
Y95{F,W}
Y97{F,W}

15 Y154{F,W}

Y158{F,W}

Y163{F,W}

Y115{F,W}

Y172{F,W}

Y175{F,W}

20 Y200{F,W}

25

30

35

Y216{F,W}

Variants with raised/lowered pI

The concept is to alter the pI for the protein such that it approaches the pH of the detergent formulation. The pI can be raised by changing negatively charged or neutral amino acids to positively charged amino acids or by changing positively charged residues to more positively charged residues. The pI can be lowered by changing positively charged or neutral amino acids to negatively charged amino acids or by changing negatively charged amino acids to more negatively charged amino acids.

Accordingly, in accordance with this embodiment the invention relates to a RP-II protease variant, in which the net electrostatic charge of the parent RP-II protease has been changed by deleting or substituting one or more negatively charged amino acid residues by neutral or positively charged amino acid residue(s), and/or by substituting one or more neutral amino acid residues by positively or negatively charged

10

15

25

30

35

amino acid residue(s), and/or by deleting or substituting one or more positively charged amino acid residue(s) by neutral or negatively charged amino acid residue(s), thereby obtaining variant which either has a lower or higher pI as compared to the pI of its parent protease.

In order to have any effect on the pI, the positions suited for substitution should be located so as to be at least partially exposed on the protein surface. It is preferred that the amino acid substitutions result in a variant protease having a pI just below the pH of the detergent.

In particular, an amino acid residue located in one or more positions of the parent RP-II protease and exposed at the surface of the molecule may be substituted:

It should be noted that, according to the invention, any one of the modifications of the amino acid sequence indicated above for the RP-II protease variants may be combined with any one of the other modifications mentioned above, where appropriate.

20 Methods of preparing RP-II proteases and variants

The novel RP-II proteases of the invention may be produced by conventional methods by fermentation of the microorganisms from which they were isolated in suitable media with subsequent purification from the fermentation broth.

However, it is preferred to use the isolated DNA sequences of the invention for the production of both the novel isolated RP-II proteases and the variants thereof. Such methods are described in detail below.

Specifically for the variants, several methods for introducing mutations into genes are known in the art. After a brief discussion of cloning RP-II protease-encoding DNA sequences (which for instance encode functional analogous of the RP-II proteases of the invention), methods for generating mutations at specific sites within the RP-II protease encoding sequence will be indicated. The mutated polynucleotide sequences are subsequently used for the production of the RP-II protease variants of the invention in a manner similar to that for producing the novel isolated RP-II proteases.

- Tame -- ___

35

The DNA sequence encoding a parent RP-II protease may be isolated from any cell or microorganism producing the RP-II protease in question by various methods, well known in the art. Useful sources producing RP-II proteases are grampositive bacteria belonging to the genus Bacillus, such as Bacillus licheniformis, Bacillus pumilus, Bacillus halmapalus, Bacillus alkalophilus, Bacillus amyloliquefaciens, Bacillus brevis. Bacillus circulans, Bacillus coagulans, Bacillus Bacillus megaterium, Bacillus Bacillus lentus. lautus, stearothermophilus. Bacillus subtilis. or Bacillus thuringiensis.

In another preferred embodiment, the nucleic acid sequences are obtained from a Bacillus licheniformis, Bacillus pumilus, or Bacillus halmapalus.

10

15

20

25

30

35

40

Strains of these species are readily accessible to the public in a number of culture collections, such as the American Type Culture Collection (ATCC), Deutsche Sammlung und Zellkulturen GmbH Mikroorganismen Centraalbureau Voor Schimmelcultures (CBS), and Agricultural Research Service Patent Culture Collection, Northern Regional Research Center (NRRL).

In another more preferred embodiment, the nucleic acid sequence is the sequence contained in plasmid pUC19/AC116, pUC/CDJ31, pUC/BO32, pUC/JA96, or pUC/AA513, which is contained in DSM 12841: E. coli pUC19/AC116, DSM 12842: E. coli pUC/CDJ31, DSM 12843: E. coli pUC/BO32, DSM 12844: E. coli pUC/JA96, and DSM 12845: E. coli pUC/AA513, respectively. In another preferred embodiment, the nucleic acid sequence is that of SEQ ID NO: 1, 3, 5, 7, 9, or 11, which encodes a mature polypeptide.

such nucleic acid sequences may Furthermore, identified and obtained from other sources including microorganisms isolated from nature (e.g., soil, composts, water, etc.) using the above-mentioned probes. Techniques for isolating microorganisms from natural habitats are well known in the art. The nucleic acid sequence may then be derived by similarly screening a genomic or cDNA library of another microorganism. _Once a nucleic acid sequence encoding a polypeptide has been _detected with the probe(s), the sequence may be isolated or cloned by utilizing techniques

which are known to those of ordinary skill in the art (see, e.g., Sambrook et al., 1989, supra).

The techniques used to isolate or clone a nucleic acid sequence encoding a polypeptide are known in the art and include isolation from genomic DNA, preparation from cDNA, or a combination thereof. The cloning of the nucleic acid sequences of the present invention from such genomic DNA can be effected, e.g., by using the well known polymerase chain reaction (PCR) or antibody screening of expression libraries to detect cloned DNA fragments with shared structural features. See, e.g., Innis et al., 1990, PCR: A Guide to Methods and Application, Academic Press, New York. nucleic acid amplification procedures such as ligase chain reaction (LCR), ligated activated transcription (LAT) and nucleic acid sequence-based amplification (NASBA) may be The nucleic acid sequence may be cloned from a strain of Bacillus, or another or related organism and thus, for example, may be an allelic or species variant of the polypeptide encoding region of the nucleic acid sequence.

10

15

20

25

30

35

...

_ 1144

More specifically, first a genomic DNA and/or cDNA library may be constructed using chromosomal DNA or messenger RNA from the organism that produces the RP-II protease to be studied. Then, if the amino acid sequence of the RP-II protease is known, homologous, labelled oligonucleotide probes may be synthesized and used to identify RP-II protease-encoding clones from a genomic library prepared from the organism in question. Alternatively, a labelled oligonucleotide probe containing sequences homologous to a known RP-II protease could be used as a probe to identify RP-II protease encoding clones, using hybridization and washing conditions of lower stringency.

Yet another method for identifying RP-II protease producing clones would involve inserting fragments of genomic DNA into an expression vector, such as a plasmid, transforming RP-II protease-negative bacteria with the resulting genomic DNA library, and then plating the transformed bacteria onto agar containing a substrate for the RP-II protease thereby allowing clones expressing the RP-II protease to be identified.

Alternatively, the DNA sequence encoding the enzyme may be prepared synthetically by established standard methods, e.g. the phosphoamidite method described by S.L. Beaucage and M.H. Caruthers, Tetrahedron Letters 22, 1981, pp. 1859-1869, or the method described by Matthes et al., The EMBO J. 3, 1984, pp. 801-805. According to the phosphoamidite method, oligonucleotides are synthesized, e.g. in an automatic DNA synthesizer, purified, annealed, ligated and cloned in appropriate vectors.

Finally, the DNA sequence may be of mixed genomic and synthetic, mixed synthetic and cDNA or mixed genomic and cDNA origin prepared by ligating fragments of synthetic, genomic or cDNA origin (as appropriate), the fragments corresponding to various parts of the entire DNA sequence, in accordance with standard techniques. The DNA sequence may also be prepared by polymerase chain reaction (PCR) using specific or degenerate primers, for instance as described in US 4,683,202 or R.K. Saiki et al., Science 239, 1988, pp. 487-491.

10

15

25

30

35

40

20 <u>Mutant Nucleic Acid Sequences and Methods for the Production</u> thereof

The present invention also relates to mutant nucleic acid sequences comprising at least one mutation in the mature polypeptide coding sequence of a functional RP-II protease analogue, and especially of SEQ ID NO:1, 3, 5, 7, 9, 11, or 13. wherein the mutant nucleic acid sequence encodes a functional analogue of a RP-II protease that may be modified in comparison to the parent protease depending on the nature of the mutation performed, especially variants of the mature polypeptide of SEQ ID NO:2, 4, 6, 8, 10, 12 or 14, or a fragment thereof which has protease activity.

Modification of a nucleic acid sequence of the present invention may be necessary for the synthesis of polypeptides substantially similar to the polypeptide. The term "substantially similar" to the polypeptide refers to non-naturally occurring forms of the polypeptide. These polypeptides may differ in some engineered way from the polypeptide isolated from its native source, e.g., variants that differ in specific activity, thermostability, phoptimum, or the like. The variant sequence may be

constructed on the basis of the nucleic acid sequence presented as the polypeptide encoding part of SEQ ID NO:1, e.g., a subsequence thereof, and/or by introduction of nucleotide substitutions which do not give rise to another amino acid sequence of the polypeptide encoded by the nucleic acid sequence, but which corresponds to the codon usage of the host organism intended for production of the enzyme, or by introduction of nucleotide substitutions which may give rise to a different amino acid sequence. For a general description of nucleotide substitution, see, e.g., Ford et al., 1991, Protein Expression and Purification 2: 95-107.

10

15

20

25

30

35

The present invention further relates to methods for producing a mutant nucleic acid sequence, comprising introducing at least one mutation into the mature polypeptide coding sequence of a functional RP-II protease analogue, and especially of SEQ ID NO:1, 3, 5, 7, 9, 11, or 13, wherein the mutant nucleic acid sequence encodes a functional analogue of a RP-II protease that may be modified in comparison to the parent protease depending on the nature of the mutation performed, especially variants of the mature polypeptide of SEQ ID NO:2, 4, 6, 8, 10, 12 or 14, or a fragment thereof which has protease activity.

The introduction of a mutation into the nucleic acid sequence to exchange one nucleotide for another nucleotide may be accomplished by site-directed mutagenesis using any of the methods known in the art. Particularly useful is the procedure which utilizes a supercoiled, double stranded DNA vector with an insert of interest and two synthetic primers containing the desired mutation. The oligonucleotide primers, each complementary to opposite strands of the vector, extend during temperature cycling by means of Pfu DNA On incorporation of the primers, a mutated polymerase. plasmid containing staggered nicks is generated. Following temperature cycling, the product is treated with DpnI which is specific for methylated and hemimethylated DNA to digest the parental DNA template and to select for mutation-_ containing synthesized DNA. Other procedures known in the art may also be used.

Once a RP-II protease encoding DNA sequence has been 40 isolated, and desirable sites for mutation identified, muta-

tions may be introduced using synthetic oligonucleotides. These oligonucleotides contain nucleotide sequences flanking the desired mutation sites; mutant nucleotides are inserted during oligonucleotide synthesis. In a specific method, a single-stranded gap of DNA, bridging the RP-II protease encoding sequence, is created in a vector carrying the RP-II protease gene. Then the synthetic nucleotide, bearing the desired mutation, is annealed to a homologous portion of the single-stranded DNA. The remaining gap is then filled in with DNA polymerase I (Klenow fragment) and the construct is ligated using T4 ligase. A specific example of this method is described in Morinaga et al., (1984, Biotechnology 2:646-639). U.S. Patent number 4,760,025, by Estell et al., issued July 1988, discloses the introduction of oligonucleotides encoding multiple mutations by performing minor alterations of the cassette, however, an even greater variety of mutations can be introduced at any one time by the Morinaga method, because a multitude of oligonucleotides, of various lengths, can be introduced.

10

15

20

25

35

Another method of introducing mutations into RP-II protease encoding sequences is described in Nelson and Long, Analytical Biochemistry 180, 1989, pp. 147-151. It involves the 3-step generation of a PCR fragment containing the desired mutation introduced by using a chemically synthesized DNA strand as one of the primers in the PCR reactions. From the PCR-generated fragment, a DNA fragment carrying the mutation may be isolated by cleavage with restriction endonucleases and reinserted into an expression plasmid.

30 Expression of RP-II proteases and variants thereof

According to the invention, a novel isolated polynucleotide or a a modified polynucleotide sequence encoding a RP-II protease or a variant thereof produced by methods described above, or any alternative methods known in the art, can be expressed, in enzyme form, using a DNA construct or an expression vector which typically includes control sequences encoding a promoter, operator, ribosome binding site, translation initiation signal, and, optionally, a repressor gene or various activator genes.

While intracellular expression may be advantageous in some respects, e.g. when using certain bacteria as host cells, it is generally preferred that the expression is extracellular. As mentioned above the RP-II proteases of the invention comprising the amino acid sequence shown in the SEQ ID NO: 1, 4, 6, 8, 10, or 12 comprise a pre-region consisting of a signal peptide and a pro-peptide permitting secretion of the expressed protease into the culture medium. If desirable, this pre-region may be substituted with a different pre-region or signal sequence, convenient accomplished by substitution of the DNA sequences encoding the respective pre-regions.

Nucleic Acid Constructs

10

15

20

25

30

35

40

The present invention also relates to nucleic acid constructs comprising a nucleic acid sequence of the present invention operably linked to one or more control sequences which direct the expression of the coding sequence in a suitable host cell under conditions compatible with the control sequences. Expression will be understood to include any step involved in the production of the polypeptide limited to, transcription, not including, but transcriptional modification, translation, post-translational modification, and secretion.

An isolated nucleic acid sequence encoding a RP-II protease of the present invention may be manipulated in a variety of ways to provide for expression of the polypeptide. Manipulation of the nucleic acid sequence prior to its insertion into a vector may be desirable or necessary depending on the expression vector. The techniques for modifying nucleic acid sequences utilizing recombinant DNA methods are well known in the art.

The term "control sequences" is defined herein to include all components which are necessary or advantageous for the expression of a RP-II protease of the present invention. Each control sequence may be native or foreign to the nucleic acid sequence encoding the polypeptide. control sequences include, but are not limited to, a leader, _ polyadenylation sequence, propeptide sequence, promoter, signal peptide sequence, and transcription terminator. minimum, the control sequences include a promoter,

transcriptional and translational stop signals. The control sequences may be provided with linkers for the purpose of introducing specific restriction sites facilitating ligation of the control sequences with the coding region of the nucleic acid sequence encoding a polypeptide. The term "operably linked" is defined herein as a configuration in which a control sequence is appropriately placed at a position relative to the coding sequence of the DNA sequence such that the control sequence directs the expression of a polypeptide.

10

15

20

25

30

35

The control sequence may be an appropriate promoter sequence, a nucleic acid sequence that is recognized by a host cell for expression of the nucleic acid sequence. The promoter sequence contains transcriptional control sequences that mediate the expression of the polypeptide. The promoter may be any nucleic acid sequence which shows transcriptional activity in the host cell of choice including mutant, truncated, and hybrid promoters, and may be obtained from genes encoding extracellular or intracellular polypeptides either homologous or heterologous to the host cell.

Examples of suitable promoters for directing transcription of the nucleic acid constructs of the present invention to produce the RP-II proteases of the invention, especially in a bacterial host cell, are the promoters obtained from the E. coli lac operon, Streptomyces coelicolor agarase gene (dagA), Bacillus subtilis levansucrase gene (sacB), Bacillus licheniformis alpha-amylase gene Bacillus stearothermophilus maltogenic amylase gene (amyM), amyloliquefaciens alpha-amylase gene (amyQ), Bacillus Bacillus licheniformis penicillinase gene (penP), Bacillus subtilis xylA and xylB genes, and prokaryotic beta-lactamase gene (Villa-Kamaroff et al., 1978, Proceedings of the National Academy of Sciences USA 75: 3727-3731), as well as the tac promoter (DeBoer et al., 1983, Proceedings of the National Academy of Sciences USA 80: 21-25). promoters are described in "Useful proteins from recombinant bacteria" in Scientific American, 1980, 242: 74-94; and in Sambrook et al., 1989, supra.

Examples of suitable promoters for directing the 40 transcription of the nucleic acid constructs of the present

<u>.</u>

invention in a filamentous fungal host cell to produce the RP-II proteases of the invention are promoters obtained from the genes for Aspergillus oryzae TAKA amylase, Rhizomucor miehei aspartic proteinase, Aspergillus niger neutral alpha-Aspergillus niger acid stable alpha-amylase, amylase, Aspergillus niger or Aspergillus awamori glucoamylase (glaA), miehei lipase, Aspergillus oryzae alkaline Rhizomucor protease, Aspergillus oryzae triose phosphate isomerase, Aspergillus nidulans acetamidase, Fusarium oxysporum trypsinlike protease (WO 96/00787), as well as the NA2-tpi promoter (a hybrid of the promoters from the genes for Aspergillus niger neutral alpha-amylase and Aspergillus oryzae triose phosphate isomerase); and mutant, truncated, and hybrid promoters thereof.

10

15

25

30

35

In a yeast host, useful promoters are obtained from the Saccharomyces cerevisiae enolase for Saccharomyces cerevisiae galactokinase (GAL1), Saccharomyces cerevisiae alcohol dehydrogenase/glyceraldehyde-3-phosphate dehydrogenase (ADH2/GAP), and Saccharomyces cerevisiae 3phosphoglycerate kinase. Other useful promoters for yeast host cells are described by Romanos et al., 1992, Yeast 8: 423-488.

also be The control sequence may transcription terminator sequence, a sequence recognized by a The terminator host cell to terminate transcription. sequence is operably linked to the 3' terminus of the nucleic acid sequence encoding the polypeptide. Any terminator that is functional in the host cell of choice may be used in the present invention.

Preferred terminators for filamentous fungal host cells are obtained from the genes for Aspergillus oryzae TAKA amylase, Aspergillus niger glucoamylase, Aspergillus nidulans anthranilate synthase, Aspergillus niger a-glucosidase, and Fusarium oxysporum trypsin-like protease.

Preferred terminators for yeast host cells are obtained for Saccharomyces cerevisiae enolase, the cytochrome C (CYC1), Saccharomyces cerevisiae glyceraldehyde-3-phosphate cerevisiae Saccharomyces dehydrogenase. - Romanos et al., 1992, supra, describe - other 40 useful terminators for yeast host cells:

The control sequence may also be a suitable leader sequence, a nontranslated region of an mRNA that is important for translation by the host cell. The leader sequence is operably linked to the 5' terminus of the nucleic acid sequence encoding the polypeptide. Any leader sequence that is functional in the host cell of choice may be used in the present invention.

Preferred leaders for filamentous fungal host cells are obtained from the genes for Aspergillus oryzae TAKA amylase and Aspergillus nidulans triose phosphate isomerase.

10

15

20

30

35

40

Suitable leaders for yeast host cells are obtained from the genes for Saccharomyces cerevisiae enolase (ENO-1), Saccharomyces cerevisiae 3-phosphoglycerate kinase, Saccharomyces cerevisiae alpha-factor, and Saccharomyces cerevisiae alcohol dehydrogenase/glyceraldehyde-3-phosphate dehydrogenase (ADH2/GAP).

The control sequence may also be a polyadenylation sequence, a sequence which is operably linked to the 3' terminus of the nucleic acid sequence and which, when transcribed, is recognized by the host cell as a signal to add polyadenosine residues to transcribed mRNA. Any polyadenylation sequence that is functional in the host cell of choice may be used in the present invention.

Preferred polyadenylation sequences for filamentous fungal host cells are obtained from the genes for Aspergillus oryzae TAKA amylase, Aspergillus niger glucoamylase, Aspergillus nidulans anthranilate synthase, Fusarium oxysporum trypsin-like protease, and Aspergillus niger alphaglucosidase.

Useful polyadenylation sequences for yeast host cells are described by Guo and Sherman, 1995, Molecular Cellular Biology 15: 5983-5990.

The control sequence may also be a signal peptide coding region that codes for an amino acid sequence linked to the amino terminus of a polypeptide and directs the encoded polypeptide into the cell's secretory pathway. The 5' end of the coding sequence of the nucleic acid sequence may inherently contain a signal peptide coding region naturally linked in translation reading frame with the segment of the coding region that rencodes the secreted polypeptide.

-- Jane Alternatively, the 5' end of the coding sequence may contain a signal peptide coding region that is foreign to the coding sequence. The foreign signal peptide coding region may be required where the coding sequence does not naturally contain a signal peptide coding region. Alternatively, the foreign signal peptide coding region may simply replace the natural signal peptide coding region in order to enhance secretion of the polypeptide. However, any signal peptide coding region, which directs the expressed polypeptide into the secretory pathway of a host cell of choice, may be used in the present invention.

10

15

20

25

30

40

_ =_

Effective signal peptide coding regions for bacterial host cells are the signal peptide coding regions obtained from the genes for Bacillus NCIB 11837maltogenic amylase, **Bacillus** stearothermophilus alpha-amylase, Bacillus licheniformis subtilisin, Bacillus betalicheniformis Bacillus stearothermophilus neutral proteases lactamase, (nprT, nprS, nprM), and Bacillus subtilis prsA. Further signal peptides are described by Simonen and Palva, 1993, Microbiological Reviews 57: 109-137.

In a preferred embodiment, the signal peptide coding region is indicated in SEQ ID NO: 1, 3, 5, 7, 9, 11, and 13, e.g. for BLC nucleotides 1 to 93 of SEQ ID NO: 1, which encodes the corresponding amino acids of SEQ ID NO: 2 4, 6, 8, 10, 12 and 14, e.g. for BLC amino acids -94 to -64 of SEQ ID NO: 2.

Effective signal peptide coding regions for filamentous fungal host cells are the signal peptide coding regions obtained from the genes for Aspergillus oryzae TAKA amylase, Aspergillus niger neutral amylase, Aspergillus niger glucoamylase, Rhizomucor miehei aspartic proteinase, Humicola insolens cellulase, and Humicola lanuginosa lipase.

Useful signal peptides for yeast host cells are obtained from the genes for Saccharomyces cerevisiae alphafactor and Saccharomyces cerevisiae invertase. Other useful signal peptide coding regions are described by Romanos et al., 1992, supra.

The control sequence may also be a propertide coding region that codes for an amino acid sequence positioned at the amino terminus of a polypeptide. The resultant

polypeptide is known as a proenzyme or propolypeptide (or a A propolypeptide is generally zymogen in some cases). inactive and can be converted to a mature active polypeptide by catalytic or autocatalytic cleavage of the propeptide from the propolypeptide. The propeptide coding region may be obtained from the genes for Bacillus subtilis alkaline protease (aprE), Bacillus subtilis neutral protease (nprT), Saccharomyces cerevisiae alpha-factor, Rhizomucor miehei aspartic proteinase, and Myceliophthora thermophila laccase (WO 95/33836).

In a preferred embodiment, the propeptide coding region is indicated in SEQ ID NO: 1, 3, 5, 7, 9, 11, and 13, e.g. for BLC nucleotides 94 to 282 of SEQ ID NO: 1 which encodes the corresponding amino acids of SEQ ID NO: 2 4, 6, 8, 10, 12 and 14, e.g. for BLC amino acids -63 to -1 of SEQ ID NO: 2.

10

15

20

25

30

35

Where both signal peptide and propeptide regions are present at the amino terminus of a polypeptide, propeptide region is positioned next to the amino terminus of the polypeptide and the signal peptide region is positioned next to the amino terminus of the propeptide region.

It may also be desirable to add regulatory sequences, which allow the regulation of the expression of polypeptide relative to the growth of the host cell. Examples of regulatory systems are those which cause the expression of the gene to be turned on or off in response to a chemical or physical stimulus, including the presence of a Regulatory systems in prokaryotic regulatory compound. systems include the lac, tac, and trp operator systems. yeast, the ADH2 system or GAL1 system may be used. filamentous fungi, the TAKA alpha-amylase Aspergillus niger glucoamylase promoter, and the Aspergillus oryzae qlucoamylase promoter may be used as regulatory Other examples of regulatory sequences are those that allow for gene amplification. In eukaryotic systems, these include the dihydrofolate reductase gene, which is amplified in the presence of methotrexate, and the metallothionein genes that are amplified with heavy metals. In these cases, the nucleic acid sequence encoding the polypeptide would be operably linked with the regulatory 40 - sequence.

It is often suitable to provide the various control or regulatory sequences from the same source.

Expression Vectors

5

10

15

20

25

30

35

The present invention also relates to recombinant expression vectors comprising a nucleic acid sequence of the invention, a promoter, and transcriptional and The various nucleic acid and translational stop signals. control sequences described above may be joined together to produce a recombinant expression vector which may include one or more convenient restriction sites to allow for insertion or substitution of the nucleic acid sequence encoding the polypeptide at such sites. Alternatively, the nucleic acid sequence of the present invention may be expressed by inserting the nucleic acid sequence or a nucleic acid construct comprising the sequence into an appropriate vector for expression. In creating the expression vector, the coding sequence is located in the vector so that the coding sequence is operably linked with the appropriate control sequences for expression.

The recombinant expression vector may be any vector (e.g., a plasmid or virus), which can be conveniently subjected to recombinant DNA procedures and can bring about the expression of the nucleic acid sequence. The choice of the vector will typically depend on the compatibility of the vector with the host cell into which the vector is to be introduced. The vectors may be linear or closed circular plasmids.

The vector may be an autonomously replicating vector, i.e., a vector that exists as an extrachromosomal entity, the which is replication of independent of chromosomal a plasmid, a bacteriophage, replication, e.g., extrachromosomal element, a minichromosome, or an artificial chromosome. The vector may contain any means for assuring self-replication. Alternatively, the vector may be one which, when introduced into the host_cell, is integrated into the genome and replicated together with the chromosome(s) _into which it has been integrated. Furthermore, a single vector or plasmid or two or more vectors or plasmids which

together contain the total DNA to be introduced into the genome of the host cell, or a transposon may be used.

The vectors of the present invention preferably contain one or more selectable markers that permit easy selection of transformed cells. A selectable marker is a gene the product of which provides for biocide or viral resistance, resistance to heavy metals, prototrophy to auxotrophs, and the like. Examples of bacterial selectable markers are the dal genes from Bacillus subtilis or Bacillus licheniformis, or markers that confer antibiotic resistance such as ampicillin, or tetracycline kanamycin, chloramphenicol resistance. Suitable markers for yeast host cells are ADE2, HIS3, LEU2, LYS2, MET3, TRP1, and URA3. A selectable marker for use in a filamentous fungal host cell may be selected from the group including, but not limited to, amdS (acetamidase), (ornithine carbamoyltransferase), bar (phosphinothricin acetyltransferase), hygB (hygromycin phosphotransferase), pyrG (orotidine-5'-phosphate niaD (nitrate reductase), decarboxylase), sC (sulfate adenyltransferase), (anthranilate synthase), as well as equivalents thereof. Preferred for use in an Aspergillus cell are the amdS and pyrG genes of Aspergillus nidulans or Aspergillus oryzae and the bar gene of Streptomyces hygroscopicus.

10

15

20

25

30

35

40

The vectors of the present invention preferably contain an element(s) that permits stable integration of the vector into the host cell genome or autonomous replication of the vector in the cell independent of the genome of the cell.

For integration into the host cell genome, the vector may rely on the nucleic acid sequence encoding the polypeptide or any other element of the vector for stable integration of the vector into the genome by homologous or non-homologous recombination. Alternatively, the vector may contain additional nucleic acid sequences for directing integration by homologous recombination into the genome of the host cell. The additional nucleic acid sequences enable the vector to be integrated into the host cell genome at a precise location(s) in the chromosome(s). To increase the likelihood of integration at a precise location, the integrational elements should preferably contain a sufficient number of inucleic acids, such as 100 to 1,500 base pairs,

preferably 400 to 1,500 base pairs, and most preferably 800 to 1,500 base pairs, which are highly homologous with the corresponding target sequence to enhance the probability of homologous recombination. The integrational elements may be any sequence that is homologous with the target sequence in the genome of the host cell. Furthermore, the integrational elements may be non-encoding or encoding nucleic acid On the other hand, the vector may be integrated sequences. of the host cell by non-homologous the genome recombination.

For autonomous replication, the vector may further comprise an origin of replication enabling the vector to replicate autonomously in the host cell in question. Examples of bacterial origins of replication are the origins of replication of plasmids pBR322, pUC19, pACYC177, and pACYC184 permitting replication in E. coli, and pUB110, pE194, pTA1060, and pAMß1 permitting replication in Bacillus. Examples of origins of replication for use in a yeast host cell are the 2-micron origin of replication, ARS1, ARS4, the combination of ARS1 and CEN3, and the combination of ARS4 and CEN6. The origin of replication may be one having a mutation which makes its functioning temperature-sensitive in the host cell (see, e.g., Ehrlich, 1978, Proceedings of the National Academy of Sciences USA 75: 1433).

More than one copy of a nucleic acid sequence of the present invention may be inserted into the host cell to increase production of the RP-II proteases of the invention. An increase in the copy number of the nucleic acid sequence can be obtained by integrating at least one additional copy of the sequence into the host cell genome or by including an amplifiable selectable marker gene with the nucleic acid sequence where cells containing amplified copies of the selectable marker gene, and thereby additional copies of the nucleic acid sequence, can be selected for by cultivating the cells in the presence of the appropriate selectable agent.

The procedures used to ligate the elements described above to construct the recombinant expression vectors of the present invention are well known to one skilled in the art (see, e.g., Sambrook et al., 1989, supra).

-

10

15

20

25

30

35

Host Cells

10

15

20

25

30

35

40

The present invention also relates to recombinant host cells, comprising a nucleic acid sequence of the invention, which are advantageously used in the recombinant production 5 of the polypeptides. A vector comprising a nucleic acid sequence of the present invention is introduced into a host cell so that the vector is maintained as a chromosomal integrant or as a self-replicating extra-chromosomal vector as described earlier. The term "host cell" encompasses any progeny of a parent cell that is not identical to the parent cell due to mutations that occur during replication. choice of a host cell will to a large extent depend upon the gene encoding the polypeptide and its source.

The host cell may be a unicellular microorganism, e.g., a prokaryote, such as a bacterial or a fungal (including yeast) cell, or a non-unicellular microorganism, e.g., a eukaryote, such as a mammal, an insect, or a plant.

Useful unicellular cells are bacterial cells such as gram positive bacteria including, but not limited to, a Bacillus cell, e.g., Bacillus alkalophilus, Bacillus amyloliquefaciens, Bacillus brevis, Bacillus circulans, Bacillus Bacillus clausii, Bacillus coaqulans, Bacillus pumilus, Bacillus halmapalus, Bacillus lentus, Bacillus licheniformis, Bacillus megaterium, Bacillus stearothermophilus, Bacillus subtilis. and Bacillus thuringiensis; or a Streptomyces cell, e.g., Streptomyces lividans or Streptomyces murinus, or gram negative bacteria such as E. coli and Pseudomonas sp. In a preferred embodiment, the bacterial host cell is a Bacillus lentus, Bacillus licheniformis, Bacillus stearothermophilus Bacillus subtilis cell. In another preferred embodiment, the Bacillus cell is an alkalophilic Bacillus.

The introduction of a vector into a bacterial host cell may, for instance, be effected by protoplast transformation (see, e.g., Chang and Cohen, 1979, Molecular General Genetics 168: 111-115), using competent cells (see, e.g., Young and Spizizin, 1961, Journal of Bacteriology 81: 823-829, or Dubnau and Davidoff-Abelson, 1971, Journal of Molecular Biology 56: 209-221), electroporation (see, e.g., Shigekawa and Dower, 1988, Biotechniques 6: 742-751), or conjugation

(see, e.g., Koehler and Thorne, 1987, Journal of Bacteriology 169: 5771-5278).

The host cell may be a eukaryote, such as a mammalian, insect, plant, or fungal cell.

5

10

15

20

25

35

40

In a preferred embodiment, the host cell is a fungal "Fungi" as used herein includes the phyla Ascomycota, Basidiomycota, Chytridiomycota, and Zygomycota (as defined by Hawksworth et al., In, Ainsworth and Bisby's Dictionary of The Fungi, 8th edition, 1995, CAB International, University Press, Cambridge, UK) as well as the Oomycota (as cited in Hawksworth et al., 1995, supra, page 171) and all mitosporic fungi (Hawksworth et al., 1995, supra).

In a more preferred embodiment, the fungal host cell is cell. "Yeast" as used herein includes yeast ascosporogenous yeast (Endomycetales), basidiosporogenous belonging to the Fungi Imperfecti and yeast (Blastomycetes). Since the classification of yeast may change in the future, for the purposes of this invention, yeast shall be defined as described in Biology and Activities of Yeast (Skinner, F.A., Passmore, S.M., and Davenport, R.R., eds., Soc. App. Bacteriol. Symposium Series No. 9, 1980).

In an even more preferred embodiment, the yeast host Kluyveromyces, Pichia, cell is а Candida, Hansenula, Saccharomyces, Schizosaccharomyces, or Yarrowia cell.

In a most preferred embodiment, the yeast host cell is a Saccharomyces carlsbergensis, Saccharomyces cerevisiae, Saccharomyces diastaticus, Saccharomyces douglasii, Saccharomyces norbensis Saccharomyces kluyveri, Saccharomyces oviformis cell. In another most preferred embodiment, the yeast host cell is a Kluyveromyces lactis 30 In another most preferred embodiment, the yeast host cell is a Yarrowia lipolytica cell.

In another more preferred embodiment, the fungal host cell is a filamentous fungal cell. "Filamentous fungi" include all filamentous forms of the subdivision Eumycota and Oomycota (as defined by_Hawksworth et al., 1995, supra). A wall composed of chitin, cellulose, mvcelial and other complex chitosan, mannan, polysaccharides characterizes the filamentous fungi. Vegetative growth is by hyphal elongation and carbon catabolism is obligately

aerobic. In contrast, vegetative growth by yeasts such as Saccharomyces cerevisiae is by budding of a unicellular thallus and carbon catabolism may be fermentative.

In an even more preferred embodiment, the filamentous fungal host cell is a cell of a species of, but not limited to, Acremonium, Aspergillus, Fusarium, Humicola, Mucor, Myceliophthora, Neurospora, Penicillium, Thielavia, Tolypocladium, or Trichoderma.

In a most preferred embodiment, the filamentous fungal host cell is an Aspergillus awamori, Aspergillus foetidus, 10 Aspergillus japonicus, Aspergillus nidulans, Aspergillus niger or Aspergillus oryzae cell. In another most preferred embodiment, the filamentous fungal host cell is a Fusarium bactridioides, Fusarium cerealis, Fusarium crookwellense, Fusarium culmorum, Fusarium graminearum, Fusarium graminum, 15 Fusarium heterosporum, Fusarium negundi, Fusarium oxysporum, Fusarium reticulatum, Fusarium roseum, Fusarium sambucinum, Fusarium sarcochroum, Fusarium sporotrichioides, Fusarium sulphureum, Fusarium torulosum, Fusarium trichothecioides, or 20 Fusarium venenatum cell. In an even most preferred embodiment, the filamentous fungal parent cell is a Fusarium venenatum (Nirenberg sp. nov.) cell. In another most preferred embodiment, the filamentous fungal host cell is a Humicola insolens or Humicola lanuginosa cell. In another most preferred embodiment, the filamentous fungal host cell is a Mucor miehei cell. In another most preferred embodiment. the filamentous fungal host cell Myceliophthora thermophila cell. In another most preferred embodiment, the filamentous fungal host cell is a Neurospora In another most preferred embodiment, the crassa cell. 30 filamentous fungal host cell is a Penicillium purpurogenum In another most preferred embodiment, the filamentous fungal host cell is a Thielavia terrestris cell. In another most preferred embodiment, the Trichoderma cell Trichoderma koningii, Trichoderma harzianum, Trichoderma 35 longibrachiatum, Trichoderma reesei or Trichoderma viride cell.

Fungal cells may be transformed by a process involving protoplast formation, transformation of the protoplasts, and 40 regeneration of the cell wall in a manner known per se.

ا اعتدار جا

Suitable procedures for transformation of Aspergillus host cells are described in EP 238 023 and Yelton et al., 1984, Proceedings of the National Academy of Sciences USA 81: 1470-1474. Suitable methods for transforming Fusarium species are described by Malardier et al., 1989, Gene 78: 147-156 and WO 96/00787. Yeast may be transformed using the procedures described by Becker and Guarente, In Abelson, J.N. and Simon, M.I., editors, Guide to Yeast Genetics and Molecular Biology, Methods in Enzymology, Volume 194, pp 182-187, Academic Press, Inc., New York; Ito et al., 1983, Journal of Bacteriology 153: 163; and Hinnen et al., 1978, Proceedings of the National Academy of Sciences USA 75: 1920.

10

15

20

25

30

35

40

The present invention therefore also relates to a transgenic plant, plant part or plant cell which has been transformed with a DNA sequence encoding the proteases or variants of the invention so as to express and produce this enzyme in recoverable quantities. The enzyme may be recovered from the plant or plant part.

The transgenic plant can be dicotyledonous or monocotyledonous, for short a dicot or a monocot. Examples of monocot plants are grasses, such as meadow grass (blue grass, Poa), forage grass such as festuca, lolium, temperate grass, such as Agrostis, and cereals, e.g. wheat, oats, rye, barley, rice, sorghum and maize (corn).

Examples of dicot plants are tobacco, legumes, such as lupins, potato, sugar beet, pea, bean and soybean, and cruciferous (family Brassicaceae), such as cauliflower, oil seed rape and the closely related model organism Arabidopsis thaliana.

Examples of plant parts are stem, callus, leaves, root, fruits, seeds, and tubers. In the present context, also specific plant tissues, such as chloroplast, apoplast, mitochondria, vacuole, peroxisomes and cytoplasm are considered to be a plant part. Furthermore, any plant cell, whatever the tissue origin, is considered to be a plant part.

Also included within the scope of the invention are the progeny of such plants, plant parts and plant cells.

The transgenic plant or plant cell expressing the enzyme of the invention may be constructed in accordance with methods known in the art. In short the plant or plant cell is

5

10

15

20

25

30

35

40

constructed by incorporating one or more expression constructs encoding the enzyme of the invention into the plant host genome and propagating the resulting modified plant or plant cell into a transgenic plant or plant cell.

Conveniently, the expression construct is a DNA construct which comprises a gene encoding the enzyme of the invention in operable association with appropriate regulatory sequences required for expression of the gene in the plant or plant part of choice. Furthermore, the expression construct may comprise a selectable marker useful for identifying host cells into which the expression construct has been integrated and DNA sequences necessary for introduction of the construct into the plant in question (the latter depends on the DNA introduction method to be used).

The choice of regulatory sequences, such as promoter and terminator sequences and optionally signal or transit sequences is determined, e.g. on the basis of when, where and how the enzyme is desired to be expressed. For instance, the expression of the gene encoding the enzyme of the invention may be constitutive or inducible, or may be developmental, stage or tissue specific, and the gene product may be targeted to a specific tissue or plant part such as seeds or leaves. Regulatory sequences are e.g. described by Tague et al, Plant, Phys., 86, 506, 1988.

For constitutive expression the 35S-CaMV promoter may be used (Franck et al., 1980. Cell 21: 285-294). Organspecific promoters may e.g. be a promoter from storage sink tissues such as seeds, potato tubers, and fruits (Edwards & Coruzzi, 1990. Annu. Rev. Genet. 24: 275-303), or from metabolic sink tissues such as meristems (Ito et al., 1994. Plant Mol. Biol. 24: 863-878), a seed specific promoter such as the glutelin, prolamin, globulin or albumin promoter from rice (Wu et al., Plant and Cell Physiology Vol. 39, No. 8 pp. 885-889 (1998)), a Vicia faba promoter from the legumin B4 and the unknown seed protein gene from Vicia faba described by Conrad U. et al, Journal of Plant Physiology Vol. 152, No. 6 pp. 708-711 (1998), a promoter from a seed oil body protein (Chen et al., Plant and cell physiology vol. 39, No. 9 pp. 935-941 (1998), the storage protein napA promoter from Brassica napus or any other seed specific promoter known in

the art, e.g. as described in WO 91/14772. Furthermore, the promoter may be a leaf specific promoter such as the rbcs promoter from rice or tomato (Kyozuka et al., Physiology Vol. 102, No. 3 pp. 991-1000 (1993), the chlorella virus adenine methyltransferase gene promoter (Mitra, A. and Higgins, DW, Plant Molecular Biology Vol. 26, No. 1 pp. 85-93 (1994), or the aldP gene promoter from rice (Kagaya et al., Molecular and General Genetics Vol. 248, No. 6 pp. 668-674 (1995), or a wound inducible promoter such as the potato pin2 promoter (Xu et al, Plant Molecular Biology Vol. 22, No. 4 pp. 573-588 (1993).

10

15

25

30

A promoter enhancer element may be used to achieve higher expression of the enzyme in the plant. For instance, the promoter enhancer element may be an intron that is placed between the promoter and the nucleotide sequence encoding the enzyme. For instance, Xu et al. op cit disclose the use of the first intron of the rice actin 1 gene to enhance expression.

The selectable marker gene and any other parts of the expression construct may be chosen from those available in 20 the art.

The DNA construct is incorporated into the plant genome according to conventional techniques known in the art, Agrobacterium-mediated transformation, virusincluding mediated transformation, micro injection, bombardment, biolistic transformation, and electroporation (Gasser et al, Science, 244, 1293; Potrykus, Bio/Techn. 8, 535, 1990; Shimamoto et al, Nature, 338, 274, 1989).

Presently, Agrobacterium tumefaciens mediated gene transfer is the method of choice for generating transgenic dicots (for review Hooykas & Schilperoort, 1992. Plant Mol. 15-38), however it can also be used for Biol. 19: transforming monocots, although other transformation methods are generally preferred for these plants. Presently, the method of choice for generating transgenic monocots is particle bombardment (microscopic gold or tungsten particles coated with the transforming DNA) of embryonic calli or developing embryos (Christou, 1992. Plant J. 2: 275-281; Shimamoto, 1994. Curr. Opin. Biotechnol. 5: 158-162; Vasil et 40 al., 1992. Bio/Technology 10: 667-674). An-alternative method

for transformation of monocots is based on protoplast transformation as described by Omirulleh S, et al., Plant Molecular biology Vol. 21, No. 3 pp. 415-428 (1993).

Following transformation, the transformants having incorporated the expression construct are selected and regenerated into whole plants according to methods well-known in the art.

Methods of Production

20

25

30

35

The present invention also relates to methods for producing a polypeptide comprising (a) cultivating a host cell under conditions suitable for production of the polypeptide; and (b) recovering the polypeptide.

The present invention also relates to methods for producing a polypeptide of the present invention comprising (a) cultivating a host cell under conditions conducive for production of the polypeptide, wherein the host cell comprises a mutant nucleic acid sequence having at least one mutation in the mature polypeptide coding region of SEQ ID NO: 1, 3, 5, 7, 9, or 11 wherein the mutant nucleic acid sequence encodes a polypeptide which consists of the amino acid sequence of the mature peptide of SEQ ID NO: 2, 4, 6, 8, 10, or 12, and (b) recovering the polypeptide.

In the production methods of the present invention, the cells are cultivated in a nutrient medium suitable for production of the polypeptide using methods known in the art. For example, the cell may be cultivated by shake flask large-scale cultivation. small-scale orfermentation (including continuous, batch, fed-batch, or solid state fermentations) industrial in laboratory or performed in a suitable medium and under conditions allowing the polypeptide to be expressed and/or isolated. cultivation takes place in a suitable nutrient medium comprising carbon and nitrogen sources and inorganic salts, using procedures known in the art. Suitable media are available from commercial suppliers or may be prepared according to published compositions (e.g., in catalogues of the American Type Culture Collection). If the polypeptide is secreted into the nutrient medium, the polypeptide can be

recovered directly from the medium. If the polypeptide is not secreted, it can be recovered from cell lysates.

The polypeptides may be detected using methods known in the art that are specific for the polypeptides. These detection methods may include use of specific antibodies, formation of an enzyme product, or disappearance of an enzyme substrate. For example, an enzyme assay may be used to determine the activity of the polypeptide as described herein.

The resulting polypeptide may be recovered by methods known in the art. For example, the polypeptide may be recovered from the nutrient medium by conventional procedures including, but not limited to, centrifugation, filtration, extraction, spray drying, evaporation, or precipitation.

The polypeptides may be purified by a variety of procedures known in the art including, but not limited to, chromatography (e.g., ion exchange, affinity, hydrophobic, size exclusion), electrophoretic chromatofocusing, and isoelectric focusing), (e.g., preparative procedures differential solubility (e.q., ammonium sulfate precipitation), SDS-PAGE, or extraction (see, e.g., Protein Purification, J.-C. Janson and Lars Ryden, editors, VCH Publishers, New York, 1989).

25 DETERGENT COMPOSITIONS COMPRISING THE ENZYMES AND VARIANTS OF THE INVENTION

The present invention comprises the use of the enzymes and variant enzymes of the invention in cleaning and detergent compositions and such compositions comprising the novel isolated RP-II proteases and RP-II protease variants or mutants. Such cleaning and detergent compositions are well described in the art and reference is made to WO 96/34946; WO 97/07202; WO 95/30011 for further description of suitable cleaning and detergent compositions.

Furthermore the example(s) below demonstrate the wash performance and improvements therein for a number of RP-II proteases and variants of the invention.

<u>Detergent Compositions</u>

15

20

30

35

The enzyme of the invention may be added to and thus become a component of a detergent composition.

The detergent composition of the invention may for example be formulated as a hand or machine laundry detergent composition including a laundry additive composition suitable for pre-treatment of stained fabrics and a rinse added fabric softener composition, or be formulated as a detergent composition for use in general household hard surface cleaning operations, or be formulated for hand or machine dishwashing operations.

10

15

20

30

35

40

specific aspect, the invention provides a In a detergent additive comprising the enzyme of the invention. The detergent additive as well as the detergent composition may comprise one or more other enzymes such as a further protease, a lipase, a cutinase, an amylase, a carbohydrase, a cellulase, a pectinase, a mannanase, an arabinase, galactanase, a xylanase, an oxidase, e.g., a laccase, and/or a peroxidase.

In general the properties of the chosen enzyme(s) should be compatible with the selected detergent, (i.e. pHoptimum, compatibility with other enzymatic and non-enzymatic ingredients, etc.), and the enzyme(s) should be present in effective amounts.

Proteases: Suitable further proteases include those 25 animal, vegetable or microbial origin. Microbial origin is preferred. Chemically modified or protein engineered mutants are included. The protease may be a serine protease or a metallo protease, preferably an alkaline microbial protease or a trypsin-like protease. Examples of alkaline proteases and subtilisins, especially those are RP-II proteases, derived from Bacillus, e.q. the RP-II proteases disclosed herein, and subtilisins, such as subtilisin Novo, subtilisin Carlsberg, subtilisin 309, subtilisin 147 and subtilisin 168 (described in WO 89/06279). Examples of trypsin-like proteases are trypsin (e.g. of porcine or bovine origin) and the Fusarium protease described in WO 89/06270 and WO 94/25583.

Examples of useful proteases are the wild type RP-II proteases and variants thereof described herein, subtilisin variants described in WO 92/19729, WO=98/20115, WO

5

10

15

98/20116, and WO 98/34946, especially the subtilisin variants with substitutions in one or more of the following positions: 27, 36, 57, 76, 87, 97, 101, 104, 120, 123, 167, 170, 194, 206, 218, 222, 224, 235 and 274.

Commercially available protease enzymes include AlcalaseTM, SavinaseTM, PrimaseTM, DuralaseTM, EsperaseTM, and KannaseTM (Novo Nordisk A/S), MaxataseTM, MaxacalTM, MaxapemTM, ProperaseTM, PurafectTM, Purafect OxPTM, FN2TM, and FN3TM (Genencor International Inc.).

It was found that special synergistic advantages could be obtained by especially combinations comprising a RP-II protease and a subtilisin of the subtilase group I-S2 (Siezen et al. Protein Science 6 (1997) 501-523) or high alkaline subtilisins. Sub-group I-S2 proteases are described as highly alkaline subtilisins and comprise enzymes such as subtilisin PB92 (MAXACAL*, Gist-Brocades NV), subtilisin 309 (SAVINASE*, NOVO NORDISK A/S), subtilisin 147 (ESPERASE*, NOVO NORDISK A/S), and alkaline elastase YaB.

The combinations of BLC and JA96 and variants thereof

with SavinaseTM and variants thereof (e.g. DuralaseTM,

KannaseTM, MaxataseTM, MaxacalTM, MaxapemTM, ProperaseTM,

PurafectTM, Purafect OxPTM, FN2TM, and FN3TM) were found to be especially useful in detergents.

Lipases: Suitable lipases include those of bacterial or fungal origin. Chemically modified or protein engineered 25 mutants are included. Examples of useful lipases include lipases from Humicola (synonym Thermomyces), e.g. from H. lanuqinosa (T. lanuqinosus) as described in EP 258 068 and EP 305 216 or from H. insolens as described in WO 96/13580, a Pseudomonas lipase, e.g. from P. alcaligenes 30 pseudoalcaligenes (EP 218 272), P. cepacia (EP 331 376), P. stutzeri (GB 1,372,034), P. fluorescens, Pseudomonas sp. 96/27002), strain SD 705 (WO 95/06720 and WO wisconsinensis (WO 96/12012), a Bacillus lipase, e.g. from B. subtilis (Dartois et al. (1993), Biochemica et Biophysica 35 Acta, 1131, 253-360), B. stearothermophilus (JP 64/744992) or B. pumilus (WO 91/16422).

Other- examples are lipase variants such as those described in WO 92/05249, WO 94/01541, EP 407 225, EP 260

105, WO 95/35381, WO 96/00292, WO 95/30744, WO 94/25578, WO 95/14783, WO 95/22615, WO 97/04079 and WO 97/07202.

Preferred commercially available lipase include Lipolase™ and Lipolase Ultra™ (Novo Nordisk A/S).

5 Amylases: Suitable amylases (α and/or β) include those of bacterial or fungal origin. Chemically modified or protein engineered mutants are included. Amylases include, example, α -amylases obtained from Bacillus, e.g. a special strain of B. licheniformis, described in more detail in GB 1,296,839.

10

15

35

Examples of useful amylases are the variants described in WO 94/02597, WO 94/18314, WO 96/23873, and WO 97/43424, especially the variants with substitutions in one or more of the following positions: 15, 23, 105, 106, 124, 128, 133, 154, 156, 181, 188, 190, 197, 202, 208, 209, 243, 264, 304, 305, 391, 408, and 444.

available amylases DuramylTM, are Commercially Termamyl[™], Fungamyl[™] and BAN[™] (Novo Nordisk A/S), Rapidase[™] and Purastar[™] (from Genencor International Inc.).

20 <u>Cellulases</u>: Suitable cellulases include those of bacterial or fungal origin. Chemically modified or protein engineered mutants are included. Suitable cellulases include cellulases from the genera Bacillus, Pseudomonas, Humicola, Fusarium, Thielavia, Acremonium, e.g. the fungal cellulases produced from Humicola insolens, Myceliophthora thermophila and 25 Fusarium oxysporum disclosed in US 4,435,307, US 5,648,263, US 5,691,178, US 5,776,757 and WO 89/09259.

Especially suitable cellulases are the alkaline or neutral cellulases having colour care benefits. Examples of 30 such cellulases are cellulases described in EP 0 495 257, EP 0 531 372, WO 96/11262, WO 96/29397, WO 98/08940. Other examples are cellulase variants such as those described in WO 94/07998, EP 0 531 315, US 5,457,046, US 5,686,593, US 5,763,254, WO 95/24471, WO 98/12307 and PCT/DK98/00299.

Commercially available cellulases include Celluzyme™, and Carezyme™ (Novo Nordisk A/S), Clazinase™, and Puradax HA™ (Genencor International Inc.), and KAC-500(B)™ (Kao _ Corporation).

Peroxidases/Oxidases: Suitable peroxidases/oxidases include those of plant, bacterial or fungal origin. Chemically modified or protein engineered mutants are included. Examples of useful peroxidases include peroxidases from Coprinus, e.g. from C. cinereus, and variants thereof as those described in WO 93/24618, WO 95/10602, and WO 98/15257.

Commercially available peroxidases include $Guardzyme^{TM}$ (Novo Nordisk A/S).

The detergent enzyme(s) may be included in a detergent composition by adding separate additives containing one or more enzymes, or by adding a combined additive comprising all of these enzymes. A detergent additive of the invention, i.e. a separate additive or a combined additive, can be formulated e.g. as a granulate, a liquid, a slurry, etc. Preferred detergent additive formulations are granulates, in particular non-dusting granulates, liquids, in particular stabilized liquids, or slurries.

10

15

20

25

35

Non-dusting granulates may be produced, disclosed in US 4,106,991 and 4,661,452 and may optionally be coated by methods known in the art. Examples of waxy coating materials are poly(ethylene oxide) products (polyethylene qlycol, PEG) with mean molar weights of 1000 to 20000; ethoxylated nonylphenols having from 16 to 50 ethylene oxide units; ethoxylated fatty alcohols in which the alcohol contains from 12 to 20 carbon atoms and in which there are 15 to 80 ethylene oxide units; fatty alcohols; fatty acids; and mono- and di- and triglycerides of fatty acids. Examples of film-forming coating materials suitable for application by fluid bed techniques are given in GB 1483591. Liquid enzyme preparations may, for instance, be stabilized by adding a polyol such as propylene glycol, a sugar or sugar alcohol, lactic acid or boric acid according to established methods. Protected enzymes may be prepared according to the method disclosed in EP 238,216.

The detergent composition of the invention may be in any convenient form, e.g., a bar, a tablet, a powder, a granule, a paste or a liquid. A liquid detergent may be aqueous, typically containing up to 70 % water and 0-30 % organic solvent, or non-aqueous.

The detergent composition comprises one or surfactants, which may be non-ionic including semi-polar and/or anionic and/or cationic and/or zwitterionic. The surfactants are typically present at a level of from 0.1% to 60% by weight.

When included therein the detergent will usually contain from about 1% to about 40% of an anionic surfactant such as linear alkylbenzenesulfonate, alpha-olefinsulfonate, alkyl sulfate (fatty alcohol sulfate), alcohol ethoxysulfate, secondary alkanesulfonate, alpha-sulfo fatty acid methyl ester, alkyl- or alkenylsuccinic acid or soap.

10

15

25

30

When included therein the detergent will usually contain from about 0.2% to about 40% of a non-ionic surfactant such as alcohol ethoxylate, nonylphenol ethoxylate, alkylpolyglycoside, alkyldimethylamineoxide, mono-ethanolamide, fatty ethoxylated fatty acid monoethanolamide, polyhydroxy alkyl fatty acid amide, or Nacyl N-alkyl derivatives of glucosamine ("glucamides").

The detergent may contain 0-65 % of a detergent builder or complexing agent such as zeolite, diphosphate, 20 phosphonate, carbonate, citrate. triphosphate, ethylenediaminetetraacetic acid, nitrilotriacetic acid, diethylenetriaminepentaacetic acid, alkyl- or alkenylsuccinic acid, soluble silicates or layered silicates (e.g. SKS-6 from Hoechst).

The detergent may comprise one or more polymers. Examples are carboxymethylcellulose, poly(vinylpyrrolidone), poly (ethylene glycol), poly(vinyl alcohol), poly-(vinylpyridine-N-oxide), poly(vinylimidazole), polycarboxylates such as polyacrylates, maleic/acrylic acid copolymers and lauryl methacrylate/acrylic acid copolymers.

The detergent may contain a bleaching system, which such as perborate, comprise H2O2 source а percarbonate, which may be combined with a peracid-forming such as tetraacetylethylenediamine activator nonanoyloxybenzenesulfonate. Alternatively, the bleaching system may comprise peroxyacids of e.g. the amide, imide, or sulfone type.

The enzyme(s) of the detergent composition of the invention may be stabilized using conventional stabilizing -4.0

agents, e.g., a polyol such as propylene glycol or glycerol, a sugar or sugar alcohol, lactic acid, boric acid, or a boric acid derivative, e.g., an aromatic borate ester, or a phenyl boronic acid derivative such as 4-formylphenyl boronic acid, and the composition may be formulated as described in e.g. WO 92/19709 and WO 92/19708.

The detergent may also contain other conventional detergent ingredients such as e.g. fabric conditioners including clays, foam boosters, suds suppressors, anticorrosion agents, soil-suspending agents, anti-soil redeposition agents, dyes, bactericides, optical brighteners, hydrotropes, tarnish inhibitors, or perfumes.

It is at present contemplated that in the detergent compositions any enzyme, in particular the enzyme of the invention, may be added in an amount corresponding to 0.01-100 mg of enzyme protein per litre of wash liquor, preferably 0.05-5 mg of enzyme protein per litre of wash liquor, in particular 0.1-1 mg of enzyme protein per litre of wash liquor.

The enzyme of the invention may additionally be incorporated in the detergent formulations disclosed in WO 97/07202, which is hereby incorporated as reference.

MATERIALS AND METHODS

25

15

STRAINS:

- B. subtilis DN1885 (Diderichsen et al., 1990).
- B. lentus 309 and 147 are specific strains of Bacillus lentus, deposited with the NCIB and accorded the accession numbers NCIB 10309 and 10147, and described in US Patent No. 3,723,250 incorporated by reference herein.
- E. coli MC 1000 (M.J. Casadaban and S.N. Cohen (1980);
 J. Mol. Biol. 138 179-207) was made r⁻,m⁺ by conventional methods and is also described in US Patent Application Serial
 No. 039,298.

PLASMIDS:

pNM1003: E. coli - B. subtilis shuttle vector, derived from pSJ3 (Described by Jacob Schiødt et al. in Protein and 5

20

30

Peptide letters 3:39-44 (1996)), containing a synthetic gene encoding for RP-II protease BLC. The construction of pNM1003 is indicated in Fig. 2.

pNM1003EXP: B. subtilis BLC expression vector.

pSX 222: B. subtilis expression vector (Described in WO 96/34946).

GENERAL MOLECULAR BIOLOGY METHODS:

Unless otherwise mentioned the DNA manipulations and
transformations were performed using standard methods of
molecular biology (Sambrook et al. (1989) Molecular cloning:
A laboratory manual, Cold Spring Harbor lab., Cold Spring
Harbor, NY; Ausubel, F. M. et al. (eds.) "Current protocols
in Molecular Biology". John Wiley and Sons, 1995; Harwood, C.
R., and Cutting, S. M. (eds.) "Molecular Biological Methods
for Bacillus". John Wiley and Sons, 1990).

ENZYMES FOR DNA MANIPULATIONS

Enzymes for DNA manipulations were used according to the specifications of the suppliers.

Unless otherwise mentioned all enzymes for DNA manipulations, such as e.g. restriction endonucleases, ligases etc., are obtained from New England Biolabs, Inc.

25 PROTEOLYTIC ACTIVITY

A GU is a Glycine Unit, defined as the proteolytic enzyme activity, which, under standard conditions, during a 15 minutes' incubation at 40° C, with N-acetyl casein as substrate, produces an amount of NH₂-group equivalent to 1 mmole of glycine.

RP-II protease activity can be measured using the PNA assay with succinyl-alanine-alanine-proline-glutamicacid-paranitroanilide as a substrate. The principle of the PNA assay is described in Rothgeb, T.M., Goodlander, B.D., Garrison, P.H., and Smith, L.A., Journal of the American Oil Chemists' Society, Vol. 65 (5) pp. 806-810 (1988).

FERMENTATION:

Fermentations for the production of the enzymes of the invention were performed at 37°C on a rotary shaking table (300 r.p.m.) in 500 ml baffled Erlenmeyer flasks containing 100 ml PS-1 medium for 5 days.

Consequently in order to make an e.g. 2 litre broth 20 Erlenmeyer flasks were fermented simultaneously.

EXAMPLE 1

10 ISOLATION OF WILD TYPE ENZYMES AND CLONING OF WILD TYPE GENES

An amino acid alignment of Glutamic Acid-specific protease (BLase) from Bacillus licheniformis ATCC 14580 (Kakudo, S., et. al J. Biol. Chem. 267:23782-23788 (1992)) and extracellular metalloprotease (mpr) (Sloma, A. et. al J.

15 Bacteriol. 172: 1024-1029 (1990)) were made. Based on the alignment the following degenerate oligo nucleotide primers coding for conserved regions have been designed for molecular screening:

20 560 sense primer:

5'- GGA TGG AGA AGC GGA AAC ACN AAY TAY GAY TAY GGN GC-3 (SEQ ID NO:18) corresponds to amino acids G-W-R-S-G-N-Y-D-Y-G (SEQ ID NO:19)

25 <u>596 sense primer:</u>

5'- CCC AAG CTT GTX GYX ACN GCN GGN CAY T-3' (SEQ ID NO:20) corresponds to amino acids V-[A/V]-T-A-G-H (SEQ ID NO:21) with a CCC and Hind III site 5' tail.

30 <u>566 antisense primer:</u>

5'- GAA TAC CGG TGA ACC GCT TTG NCM NCC RTA NGT RTC-3'
(SEQ ID NO:22)
corresponds to amino acids D-T-Y-G-[G/C/W/end]-Q-S-G-S-P-V-F
(SEQ ID NO:23)

35 594 antisense primer:

5'-GCT CTA GAG TYD ATN GCN CCR TAR TC-3' (SEQ ID NO:24) corresponds to amino acids D-Y-G-A-I-[E/K] (SEQ ID NO:25) with a GC and Xba I site 5' tail.

65

where N = A, C, G or T; R = A or G; Y = C or T; D = A, G or T; X = deoxyinosine.

The genomic DNA from Bacillus strain AC116 and Bacillus strain CDJ31 were isolated according to the following procedure:

Procedure for isolating genomic DNA.

- 1. Harvest 1.5 ml culture and resuspend in 100 μ l TEL. Leave at 37C for 30 min.
 - 2.Add 500 μ l thiocynate buffer and leave at room temperature for 10 min.
 - 3.Add 250 µl NH4Ac and leave at ice for 10 min.
 - 4.Add 500 μ l CIA and mix.
- 15 5.Transfer to a microcentrifuge and spin for 10 min. at full speed.
 - 6.Transfer supernatant to a new Eppendorf tube and add 0.54 volume cold isopropanol. Mix thoroughly.
 - 7.Spin and wash the DNA pellet with 70 % EtOH.
- 20 8. Resuspend the genomic DNA in 100 μ l TER.

TE 10 mM Tris-HCl, pH 7.4

1 mM EDTA, pH 8.0

TEL 50 mg/ml Lysozym in TE-buffer

Thiocyanate 5M guanidium thiocyanate

100 mM EDTA

0.6 % w/v N-laurylsarcosine, sodium salt.

60 g thiocyanate, 20 ml 0.5 M EDTA, pH 8.0,

20 ml H₂O dissolves at 65C. Cool down to RT

and add 0.6 g N-laurylsarcosine. Add H2O to

100 ml and filter it through a 0.2 μ

sterile filter.

NH₄Ac 7.5 M CH₃COONH₄

TER 1 μ g/ml Rnase A in TE-buffer

CIA Chloroform/isoamyl alcohol 24:1

Experimental procedure

Approximately 100 to 200 ng genomic DNA is used as template for PCR amplification in PCR buffer (10 mM Tris-HCl,

pH 8.3, 50 mM KCl) containing 200 μ M of each dNTP, 3.5 mM MgCl2, 2.5 Units AmpliTaq GoldTM, and 100 pmol of each of the degenerate primers 594 and 596 or each of the degenerate primers 560 and 566. The total volume is 50 μ l. The PCR reaction is carried out in a Perkin - Elmer GeneAmp PCR System 2400. The PCR reaction is performed using a cycle profile of:

94°C - 10 min; 1 cycle

25

30

35

94°C - 1 min, 60°C - 1 min, 72°C - 30 sec; 2 cycles

10 94°C - 1 min, 59°C - 1 min, 72°C - 30 sec; 2 cycles

94°C - 1 min, 58°C - 1 min, 72°C - 30 sec; 2 cycles

15 94°C - 1 min, 52°C - 1 min, 72°C - 30 sec; 2 cycles 94°C - 1 min, 50°C - 1 min, 72°C - 30 sec; 14 cycles 72°C - 7 min; 1 cycles

5 μ l aliquots of the amplification products are analysed by electrophoresis in 1.5% agarose gels.

Purification and sequencing of PCR bands

The PCR fragments can be purified and sequenced using GFXä PCR DNA and Gel Band Purification Kit (Pharmacia Biotech) according to the manufacturer's instructions. The nucleotide sequences of the amplified PCR fragments are determined directly on the purified PCR products using 200-300 ng as template, the Taq deoxy-terminal cycle sequencing kit (Perkin-Elmer, USA), fluorescent labelled terminators and 5 pmol of either sense or antisense primer on an ABI PRISM 377 DNA Sequencer, Perkin Elmer.

PCR fragments were generated on genomic DNA from Bacillus strain AC116 and Bacillus strain CDJ31 with primer set 594/596 and primer set 560/566, purified and sequenced as described above and the DNA sequence were deduced.

Sequence between primer 596 and 594 in the 596/594 PCR fragment from Bacillus strain AC116 (5'- to 3'-):

GCGTCTATGACACGCCAAGCCGATCATTCGCGGGAACCGCCACCGTTTCCCCGGGACGAAA CGGTTCAGCTTACCGTTACGGATCTGTTACATCGACCCGCTATTTCATCCCGTCGGGTTGG CAGAGCGGAAATTCCAATTAT (SEO ID NO:26)

and translated into amino acid sequence:

5 CVYDTASRSFAGTATVSPGRNGSAYPYGSVTSTRYFIPSGWOSGNSNY (SEQ ID NO:27)

Sequence between primer 560 and 566 in the 560/566 PCR fragment from Bacillus strain AC116 (5'- to 3'-):

10

GATCGAGCTCAGCCAGCCGATCGGCAATACCGTCGGATATTTCGGATATTCATACACCGCT TCATCGCTTGCAGGAGCAGCGTGACCATCAGCGGATATCCAGGAGACAAAACAACAGGCA CCCAGTGGCAAATGTCCGGAACGATCGCTGTTTCAGAAACGTATAAACTGCAATATGCGAT C (SEQ ID NO:28)

and translated into amino acid sequence: 15 IELSOPIGNTVGYFGYSYTASSLAGAGVTISGYPGDKTTGTOWOMSGTIAVSETYKLQYAI (SEQ ID NO:29)

Sequence between primer 596 and 594 in the 596/594 PCR 20 fragment from Bacillus strain CDJ31 (5'- to 3'-):

GCATTTATGACACAGCGAGCGGGTCATTCGCCGGAACCGCTACCGTTTCTCCGGGACGGAA CGGTTCAACATATCCGTACGGATCAGTTACATCAACCCGCTATTTCATCCCGTCAGGCTAT CGAAGCGGAAATTCGAATTAC (SEQ ID NO:30)

and translated into amino acid sequence: 25 CIYDTASGSFAGTATVSPGRNGSTYPYGSVTSTRYFIPSGYRSGNSNY (SEO ID NO:31)

Sequence between primer 560 and 566 in the 560/566 PCR 30 fragment from Bacillus strain CDJ31 (5'- to 3'-):

CATAGAGCTCAGCCAGCCGATCGGCAACACCGTCGGGTATTTCGGATATTCCTACACCACC TCGTCTCTCGTTGGGTCAAGCGTTACCATCATCGGATATCCAGGCGACAAAACATCGGGCA CCCAATGCCAGATGTCCGGAAATATCGCCGTCTCAGAAACATATAAACTGCAATATGCGAT

35 C (SEQ ID NO:32) and translated into amino acid sequence:

IELSQPIGNTVGYFGYSYTTSSLVGSSVTIIGYPGDKTSGTQWQMSGNIAVSETYKLQYAI (SEQ ID NO:33) __

40 Cloning by inverse PCR:

Based on the above DNA sequences oligo nucleotide primers were designed for inverse PCR.

Bacillus strain AC116:

5 602: 5'- CGT AAG GGT AAG CTG AAC C -3' (SEQ ID NO:34) 603: 5'- CAG GAG ACA AAA CAA CAG CAG GC - 3' (SEQ ID NO:35)

Bacillus strain CDJ31:

598: 5'-GTC CCG GAG AAA CGG TAG -3' (SEQ ID NO:36)

10 600: 5'- CAC CAC CTC GTC TCT CGT TG-5' (SEQ ID NO:37)

Method for inverse PCR:

- 1.Digested 0.5-1.0 mg genome DNA with BamHI, HindIII, KpnI, PstI, XbaI and XhoI respective in a volume of 50ml over night at 37C.
 - 2. Purify the six DNA digests over a GFX Column according to manufactures instructions (GFX PCR DNA and Gel Band Purification Kit, Pharmacia Biotech).
- 3.Diluted to a final concentration of 1-10 mg/ml in ligase buffer. Add T4 ligase and incubated over night at 16C.
 4.Set up PCR as described with a long range PCR system.

PCR condition for Expand Long Template PCR System from Boehringer Mannheim with a suspected fragment length at 4-6 kb:

1 ml of ligation mixture (template)
50 pmol of each primer (Tm should be between 63C and 68C)
1 ml 20mM dNTP

5 ml 10 X Buffer 1 with MgCl2
0.75 ml Expand DNA polymerase mix (Taq and Pwo)
H2O to 50 ml

Cycle profile: 1x (94C for 2 min.)

10x (94C for 10s; 60C (depending of primer Tm) for 30s; 68C

for 4 min.)

20x (94C for 10s; 60C for 30s; 68C for 4 min. + 20s

additional added per cycle)

1x (68C for 7min.)

15

25

Gel purify the PCR products of interest (GFX) and the sequence of the gene can be determent. Based on the sequences new PCR primers for amplification and cloning of the gene can be design.

The same method were used for isolation of S2b proteases from Bacillus strain JA96, Bacillus strain BO32 and Bacillus strain AA513 with few modifications. New primers were design for molecular screening based on a new amino acid alignment containing the amino acid sequence from AC116 and CDJ31 (primer 611) and based on the N-terminal amino acid determination of the S2b protease from Bacillus strain C3371 (primer 646):

611 antisense primer:

15 5'- GCT CTA GAC GTY TTR TCX CMX GGR WAN CC-3 (SEQ ID NO:38)

corresponds to amino acids G-[Y/F]-P-[G/C]-D-K-T (SEQ ID NO:39) with a GC and XbaI site 5' tail.

20 <u>646 sense primer:</u>

5'- CCC AAG CTT GTX GTX ATH GGX GAY GAY GG-3 (SEQ ID NO:40) corresponds to amino acids V-V-I-G-G-D-D-G (SEQ ID NO:41) with a CCC and Hind III site 5' tail.

25

40

5

10

The 611/646 primer set was used as described above on genomic DNA from Bacillus strain JA96, Bacillus strain BO32 and Bacillus strain AA513 isolated as described above. Sequence determination of the PCR fragments, design of primers for inverse PCR, inverse PCR reactions and sequencing of the genes were done as described above.

All five genes were cloned into the pUC19 vector and transformed into the *Escherichia coli* strain DH10B (Life Technologies) and deposited with DSM as indicated above.

35 DSM 12841: E. coli pUC19/AC116, DSM 12842: E. coli pUC/CDJ31 DSM 12843: E. coli pUC/BO32, DSM 12844: E. coli pUC/JA96 DSM 12845: E. coli pUC/AA513

The DNA sequences and the amino acid sequences derived therefrom are indicated in SEQ ID. NOS. 1 to 12.

The coding-region of the genes can be excised from the

pUC19 constructions and subcloned into a Bacillus expression vector such as pNM1003exp, and further transformed into B. subtilis DN1885 for expression of the novel isolated RP-II proteases of the invention as described in Example 2.

5

EXAMPLE 2

CONSTRUCTION AND EXPRESSION OF ENZYME VARIANTS:

A B. subtilis-E. coli shuttle vector, pNM1003, suited to a gene coding for RP-II protease BLC and its mutants was constructed. It is derived from the B. subtilis expression vector pSX222 (Described in WO 96/34946) according to flowchart shown in Figs. 2a and 2b and as described below.

15 pKH400:

20

A DNA fragment from pJS3 encoding the beta-lactamase gene and oriC was prepared by PCR using primers introducing BamHI sites in the fragment terminals. The PCR product was digested with BamHI and ligated with BamHI digested pSX222. The ligation mixture was used to transform competent E. coli MC1000 r⁻ m⁺, selecting for ampicillin resistance.

pKH401:

A PstI site was introduced by site directed

mutagenesis in the upstream region of the gene encoding
Savinase.

pCC1:

pKH401 was restricted with PstI and MluI in order to

remove the gene encoding subtilisin 309. A 5174 bp PstI-MluI

pKH401 vector fragment was ligated with a PstI-MluI DNA

fragment encoding BLC. Such DNA can be obtained in a manner

as described in EP 482 879. The ligation mixture were used to

transform competent E. coli MC1000 r m. A plasmid (pCC1)

with a single nucleotide deletion in the BLC gene was

isolated since expression of BLC in E. coli is toxic. The

single nucleotide deletion was located in the pro-peptide

region of BLC.

71

A PmeI-BstEII fragment of pCC1 was replaced by a 343 bp RsaI-BstEII fragment from a wt BLC gene. The ligation mixture was used to transform competent B. subtilis DN1885 selecting for protease activity. Plasmid DNA was isolated and 5 verified by sequencing.

Plasmid pNM1001:

10

15

25

30

35

pNM1000 were restricted with BamHI and the 4350 bp large fragment was isolated. The 4350 bp fragment was ligated and the ligation mixture was used to transform competent B. subtilis DN1885 selecting for protease activity. Plasmid DNA was isolated and verified by DNA sequencing.

Plasmid pNM1002:

A PCR product covering the region PstI-BstEII of pNM1001, introducing a HindIII site by site-directed mutagenesis in the propeptide region BLC gene, was restricted with PstI-BstEII and ligated to a 3925 bp PstI-BstEII fragment of pNM1001. The ligation mixture was used to 20 transform competent B. subtilis DN1885 selecting for protease activity. Plasmid DNA was isolated and verified by DNA sequencing.

Plasmid pNM1003:

The ampicillin gene and oriC region of pNM1000 were amplified by PCR using primers introducing HindIII sites in the terminals. The PCR product was restricted with HindIII and ligated to HindIII restricted pNM1002. The ligation mixture was used to transform competent E. coli MC1000 r m', selecting for ampicillin resistance. Plasmid DNA was isolated and confirmed by sequencing.

pNM1003EXP:

pNM1003 was restricted with HindIII and a 4350 bp DNA fragment was isolated and ligated. The ligation mixture were used to transform competent B. subtilis DN1885, selecting for protease activity.

SITE-DIRECTED MUTAGENESIS:

BLC site-directed variants of the invention comprising specific substitutions, insertions or deletions in the molecule were made by traditional cloning of DNA fragments (Sambrook et al., Molecular Cloning: A Laboratory Manual, 2nd Ed., Cold Spring Harbor, 1989) produced by PCR of oligos containing the desired insertions (see below).

The template plasmid DNA was pNM1003, or an analogue of this containing a variant of RP-II protease BLC.

Insertions were introduced by oligo-directed mutagenesis to the construction of substitution, insertion or deletion variants resulting in RP-II BLC variants.

10

15

30

35

The BLC variants were transformed into *E. coli*. DNA purified from a over night culture of these transformants were transformed into *B. subtilis* by restriction endonuclease digestion, purification of DNA fragments, ligation, transformation of *B. subtilis*. Transformation of *B. subtilis* was performed as described by Dubnau et al., 1971, J. Mol. Biol. 56, pp. 209-221.

20 LOCALIZED RANDOM MUTAGENESIS IN ORDER TO INSERT RANDOM INSERTIONS IN A LOCALIZED REGION:

The overall strategy used to perform localized random mutagenesis was:

A mutagenic primer (oligonucleotide) was synthesized, that corresponds to the DNA sequence flanking the site of substitution, insertion or deletion, separated by the DNA base pairs defining the substitution, insertion or deletion.

Subsequently, the resulting mutagenic primer was used in a PCR reaction with a suitable opposite primer. The resulting PCR fragment was purified and extended in a second PCR-reaction, before being digested by endonucleases and cloned into the *E. coli - B. subtilis* shuttle vector (see below).

Alternatively, and if necessary, the resulting PCR fragment is used in a second PCR reaction as a primer with a second suitable opposite primer to allow digestion and cloning of the mutagenized region into the shuttle vector. The PCR reactions are performed under normal conditions.

Following this strategy a localized random library was constructed in BLC wherein substitutions were introduced at position 36.

The mutations were introduced by mutagenic primers (see below), so that all 20 amino acids, except Trp and Met, are represented (N = 25% of A, T, C, and G; whereas H = 33% A, 33% C and 33% T. The produced PCR fragment were extended towards the N-terminal of BLC by another round of PCR by combination of a overlapping sequence with a PCR-fragment produced by PCR-amplification with primers; 5' GCA CGG ACC GTT GCA GTT CGT TCT GGA GC 3' (sense) (SEQ ID NO:42) and 5' CCG GCA AAG TGA ATG AAA CAA AGG AAA AAG CGG 3' (anti-sense) (SEQ ID NO:43). The extended DNA-fragments were cloned into the BstE II- and PinA I- sites of the modified plasmid pNM1003 (see above), and ten randomly chosen E. coli colonies were sequenced to confirm the mutations designed.

10

15

20

25

30

35

The mutagenic primer (5' A TGC ACC GGA TGG NNH ATA GGT CCG AAA ACC 3'(anti-sense) (SEQ ID NO:44)) were used in a PCR reaction with a suitable sense opposite primer, situated downstream of the MluI site in pNM1003 (e.g. 5'- CCC TTT AAC CGC ACA GCG TTT -3' (anti-sense)) (SEQ ID NO:45) and the plasmid pNM1003 as template. This resulting PCR product was cloned into the pNM1003 shuttle vector by using the restriction enzymes BstE II and PinA I.

The random library was transformed into *E. coli* by well-known techniques. The library prepared contained approximately 100,000 individual clones/library. Ten randomly chosen colonies were sequenced to confirm the mutations designed.

In order to purify a BLC variant of the invention, the pNM1003EXP plasmid comprising a variant of the invention was created by digestion of pNM1003 with HindIII, ligated and transformed into a competent B. subtilis strain, selecting for protease activity, and was fermented as described above in a medium containing 10 $\mu g/ml$ Chloramphenicol (CAM).

EXAMPLE 3 PURIFICATION OF ENZYMES AND VARIANTS

This procedure relates to purification from fermentation in 2 litres scale for the production of the proteases of the invention in a *Bacillus* host cell.

Approximately 1.6 litres of fermentation broth were centrifuged at 5000 rpm for 35 minutes in 1-litre beakers. The supernatants were adjusted to pH 7 using 10% acetic acid and filtered through a Seitz Supra S100 filter plate.

At room temperature, the filtrate was applied to a 100 ml Bacitracin affinity column equilibrated with 0.01 M dimethylglutaric acid, 0.1 M boric acid and 0.002 M calcium chloride adjusted to pH 7 with sodium hydroxide (Buffer A). After washing the column with Buffer A to remove unbound protein, the protease was eluted from the Bacitracin column using Buffer A supplemented with 25% 2-propanol and 1 M sodium chloride.

10

15

20

25

30

35

The fractions with proteolytic activity from the Bacitracin purification step were combined and applied to a 750 ml Sephadex G25 column (5 cm dia.) equilibrated with Buffer A.

Fractions with proteolytic activity from the Sephadex G25 column were combined and the pH was adjusted to pH 6 with 10% acetic acid and applied to a 150 ml CM Sepharose CL 6B cation exchange column (5 cm dia.) equilibrated with a buffer containing 0.01 M dimethylglutaric acid, 0.1 M boric acid, and 0.002 M calcium chloride adjusted to pH 6 with sodium hydroxide.

The protease was eluted using a linear gradient of 0-0.2 M sodium chloride in 2 litres of the same buffer.

Finally, the protease containing fractions from the CM Sepharose column were combined and filtered through a 0.2 μm filter.

By using the techniques of Example 1 for the isolation of wild type enzymes, and the above isolation procedure the RP-II proteases indicated below were produced and isolated.

For ease of reference the wild type RP-II proteases have in Figs 1a to 1c been aligned to the RP-II protease from Bacillus licheniformis, BLC, in the manner described above to establish the numbering of the amino acid residues.

By using the techniques of Example 2 for the 40 construction of variants and fermentation, and the above

isolation procedure the following RP-II protease variants of the BLC protease were produced and isolated:

V1F

5 S5P+Y95F+E152N+Y154L

D6A

D7A

D7G+T125S+E152G+N182I

T9R

10 T15R

Y19F

H24Q+E152G+E173G

S28R; S28R+T80K; S28R+T80K+Q157R; S28R+E152V;

S28R+E152R+Y154F+Q157A; S28R+E152A+E209A; S28R+Q157R

15 I29A; I29T; I29S; I29P; I29A+E152A; I29T+E152A; I29S+E152A; I29A+E152A+E209A

C32A

M36A; M36S; M36V+M16OS; M36A+M16OS; M36L+M16OS; M36I+M16OS; M36T+M16OS

20 I37V+O143R+E152A

G46A+Y50F+D51S+E152G

H47A

C48A

Y50F

25 D51K; D51R; D51A; D51N+V77I+T137R+H144R; D51A+E152A

S53T+Y200F+E209K

S54R+E152G+Y154F

G55R

G59R+I150T

30 T62S+E152K; T62S+E152G+E209Q; T62S+E152G+Q174R+T179S

T62S+E152G+Q174R+T179S

G69R; G69L+S71A+E152A; G69R+S76T; G69R+E101A+E152A+E173A;

G69R+E152A; G69R+E152G; G69R+E152V

T70I+E152G+T179S

35 S76H

T80K; T80K+Q157R; T80K+E152A+E209A

Y82F

S90K+E152G

N94K+E152A

40 Y95F; Y95F+I129V+E152R+Y154T+Q157L; Y95F+E152R+Y154S;

Y95F+E152R+Y154S+Q157S; Y95F+E152R+Y154S+Q157F; Y95F+E152R+Y154T+Q157H; Y95F+E209K D96A E101A; E101A+E152G+E173A; E101A+E152A+E173A+E209A; 5 E101A+E173A+E209A L102V+E104R+E152A E104A; E104K; E104R; E104A+E152G; E104A+E152A; E104A+E152A+E209A; E104A+E152A+V189I; E104K+Q174R+S186A T109R 10 S116R S119A+S144T+Q157R+S175I T128K Y132F D135A 15 H141A S145P+E152G+Q157R, I150T+E152G E152K; E152R; E152A; E152V; E152G; E152A+E104A; E152A+E104A+E209A; E152A+E104A+V189I; E152A+E104A+V189L; 20 E152A+Q122R; E152R+Y154S+Q157S; E152R+Y154F+Q157S; E152R+Y154A+Q157L; E152R+Y154L+Q157Y; E152S+Y154F+Q157L; E152A+Q157R+E209A; E152K+A159S+E173D; E152G+G164R; E152G+G164R; E152A+F172Y+E173Q+Q174E; E152A+E173A; E152A+E173Q+Q174P; E152G+E173I+Q174V; E152G+N180S+T191S; 25 E152A+V189I; E152A+V189L; E152G+S199P; E152V+Y200H; E152A+E209A; E152A+D212A; E152G+D212N Y154F+Q157R Q157R Y158F 30 M160A; M160S D161A T162M; T162K; T162R; T162A; T162S S167A E173A; E173K; E173R 35 Q174R C181A N182T+C185A C185A V189I; V189L

40 H190A; H190T; H190S; H190P; H190M; H190K; H190R-

T191S; T191V; T191G; T191I; T191R+G196C

N192*

Y195F

T207R

5 E209A; E209K; E209R

D212A; D212K; D212R

Similarly the protease variants indicated below were produced and isolated from the RP-II protease from JA96

10

E155A

E155G

E155A+D156A

D156A

15 D156G

These variants exhibited better wash performance than the RP-II protease BLC in a preliminary assay.

20 EXAMPLE 4

WASH PERFORMANCE OF PROTEASE VARIANTS (I)

The following examples provide results from a number of washing tests that were conducted under the conditions indicated below.

25

EXPERIMENTAL CONDITIONS

Detergent	OMO color, 4.0 g/l
рН	10.25
Water hardness	18°dH ~ 3.22 mM Ca²+/Mg²+
Wash time	20 min.
Temperature	30°C
Enzyme conc.	10 nM
Test system	150 ml beakers with a stirring rod
Test material	5 pieces of test material (Ø 2.5 cm) in 50 ml detergent solution

Table 1. Experimental set-up.

Water hardness was adjusted by adding CaCl₂ and MgCl₂ to deionized water.

5

DETERGENT

The detergent used was obtained from a supermarket in Bagsvaerd, Denmark. Prior to use all enzymatic activity in the detergent was inactivated by microwave treatment.

10

TEST MATERIALS

The test material used was EMPAl16 (obtained from EMPA Test materials, Movenstrasse 12, CH-9015 St. Gallen, Switzerland), and cotton soiled with grass juice.

15

25

REFLECTANCE

Reflectance measurements of the test materials were done at 460 nm using a J&M Tidas MMS/16 photometer equipped with a CLX 75W Xenon lamp and fibre optics. Each textile piece was measured individually with other textile pieces (same settings) as background.

EVALUATION

The evaluation of the wash performance of the RP-II proteases was performed by measuring the reflectance of test material washed with the RP-II proteases. High reflectance

values mean that the test material was cleaned, and indicate an improved RP-II protease wash performance.

SAS 6.12 software was used to make an analysis of variance and a t-test comparison (Student-Newman-Keuls) at 95% significance on the experimental data.

RESULTS

The capital letters designate statistical groupings within each column based on a t-test. If two variants are in the same group (same letter), they cannot be separated statistically.

Enzyme	EMPA116	Grass
E152A	24.5 A	51.8 C
E152A+E209A	24.5 A	53.4 A
E152A+I29T	24.5 A	51.0 D
E152A+V144L	24.2 A	52.7 B
E152A+I29S	24.1 B	51.1 D
E104A	23.8 B	51.1 D
E152A+I29A	23.6 C	50.3 D
E173A	23.4 D	50.2 E
E209A	23.3 D	51.5 C
D212A	22.4 E	49.9 E
V189I	22.1 E	48.0 F
BLC	21.9 E	47.3 G
Blind	19.4 F	45.9 G
Root MSE	0.4	0.5
R-square	0.96	0.97

Table 2. Mean reflectance and category for each variant.

EXAMPLE 5

15

- WASH PERFORMANCE OF PROTEASE VARIANTS (II)

EXPERIMENTAL CONDITIONS

The washing tests were conducted under the same experimental conditions as described in Example 4.

5 EVALUATION

Evaluation of the RP-II proteases was done as in Example 4, except that no statistical analysis was carried out.

10 RESULTS

The reflectance measurements are shown in table 3 to 7 below.

Enzyme	EMPA116	Grass
E104K	24.2	50.7
T62S+E152G	24.2	50.5
E104K+Q204R	23.7	50.6
E209R	23.7	50.2
Y154K+Q157R	22.5	46.4
T80K+Q157R	22.4	47.8
BLC	21.9	47.3
Blind	19.4	45.9

Table 3. Mean reflectance for each variant.

Enzyme	OMO	color	/EMPA 1	16
	5 nM		10	nM
L102V+E104R+E152A	26,85	A	27,36	B/C
G69R+E152A	26,17	В	28,57	A
G69R+E152G	25,97	В	27,88	В
E152R+Y154L+Q157Y	25,13	С	27,18	B/C
Y95F+E152R+Y154S+Q157F	25,1	С	26,55	C/D
Y95F+E152R+Y154T+Q157H	24,74	С	26,7	C/D
Y95F+E152R+Y154S	24,68	С	26,69	C/D
Y95F+I129V+E152R+Y154T+Q157L	25,37	С	26	D
N94K+E152A	24,95	С	26,08	D
BLC -	23,05	D	24,53	E

Table 4. Mean reflectance and category for each variant.

Enzyme	OMO	regular/EMPA	116
	5 Nm	1	0 nM
S54R+E152G+Y154F	14,66 A	16,14	A
T70I+E152G+T179S	14,47 A	/B 16	A
T62S+E152G+E209Q	14,26 B	15,01	В
E152G+S199P	14,23 B	14,61	C
D156G*	13,95 C	13,76	E/F
E155A*	13,92 C	/D 14,13	D
JA96*	13,73 C	/D/E 14,06	D
E155A+D156A*	13,72 C	/D/E 14,24	D
E155G*	13,62 C	/D/E 14,19	D
BLC	13,5 I	E 13,83	Е

*RP-II protease JA96 and variants thereof

Table 5. Mean reflectance and category for each variant.

Enzyme	OMC	colo	or/EMPA 1	16
	5 nM		10	nM
T70I+E152G+T179S	27,39	A	28,35	A
S54R+E152G+Y154F	26,32	В	27,76	В
E152G+S199P	26,3	В	27,48	В
T62S+E152G+E209Q	26,07	В	26,56	С
BLC	24,01	С	25,24	D

Table 6. Mean reflectance and category for each variant.

Enzyme	OMO C	colo	r/EMPA116	OMO Regular/EMPA116		
	5nM/:	L	10nM/L	5nM/L	10nM/L	
G69R	25,65	A	25,55 A	13,42 A	13,56 A	
E152G+G164R	24,68	В	25,93 A	13,50 A	13,79 A	
Y82F+Y95F	24,08	В	24,50 B	13,08 B	13,55 A	
Y19F	23,06	С	23,71 C	13,07 B	13,28 A	
Y195F	22,84	С	23,45 C	12,43 C	13,27 A	
BLC	22,65	С	23,12 C	13,13 B	13,35 A	

Table 7. Mean reflectance and category for each variant.

_

<u>fil</u>

..___

PCT

Original (for SUBMISSION) - printed on 30.08.2000 01:21:07 PM

0-1	Form - PCT/RO/134 (EASY) Indications Relating to Deposited Microorganism(s) or Other Biological Material (PCT Rule 13bls)	
0-1-1	Prepared using	PCT-EASY Version 2.91
		(updated 01.07.2000)
0-2	International Application No.	PCT/DK 00/00476
0-3	Applicant's or agent's file reference	5665.204-WO
1	The indications made below relate to the deposited microorganism(s) or other biological material referred to in the description on:	
1-1	page	5
1-2	line	6
1-3	Identification of Deposit	
1-3-1	Name of depositary institution	DSMZ-Deutsche Sammlung von Mikroorganismen und Zellkulturen GmbH
1-3-2	Address of depositary institution	Mascheroder Weg 1b, D-38124
1-0-2	Address of depositing institution	Braunschweig, Germany
1-3-3	Date of deposit	04 June 1999 (04.06.1999)
1-3-4	Accession Number	DSMZ 12841
1-4	Additional Indications	NONE
1-5	Designated States for Which	all designated States
	Indications are Made	all designated States
1-6	Separate Furnishing of Indications These indications will be submitted to	NONE
	the International Bureau later	
2	The indications made below relate to the deposited microorganism(s) or other biological material referred to in the description on:	
2-1	page	5
2-2	line	6
2-3	Identification of Deposit	
2-3-1	Name of depositary institution	DSMZ-Deutsche Sammlung von
		Mikroorganismen und Zellkulturen GmbH
2-3-2	Address of depositary institution	Mascheroder Weg 1b, D-38124
		Braunschweig, Germany
2-3-3	Date of deposit	04 June 1999 (04.06.1999)
2-3-4	Accession Number	DSMZ 12842
2-4	Additional Indications	NONE
2-5	Designated States for Which Indications are Made	all designated States
2-6	Separate Furnishing of Indications	NONE
	These indications will be submitted to the International Bureau later	-

PCT

Original (for SUBMISSION) - printed on 30.08.2000 01:21:07 PM

3	The indications made below relate to the deposited microorganism(s) or	
	other biological material referred to in the description on:	
3-1	page	5
3-2	line	6
3-3	Identification of Deposit	
3-3-1	Name of depositary institution	DSMZ-Deutsche Sammlung von
		Mikroorganismen und Zellkulturen GmbH
3-3-2	Address of depositary institution	Mascheroder Weg 1b, D-38124
		Braunschweig, Germany
3-3-3	Date of deposit	04 June 1999 (04.06.1999)
3-3-4	Accession Number	DSMZ 12843
3-4	Additional Indications	NONE
3-5	Designated States for Which Indications are Made	all designated States
3-6	Separate Furnishing of Indications	NONE
	These indications will be submitted to the International Bureau later	
4	The Indications made below relate to	
	the deposited microorganism(s) or other biological material referred to in the description on:	
4-1	page	5
4-2	line	6
4-3	Identification of Deposit	
4-3-1	Name of depositary institution	DSMZ-Deutsche Sammlung von
		Mikroorganismen und Zellkulturen GmbH
4-3-2	Address of depositary institution	Mascheroder Weg 1b, D-38124
		Braunschweig, Germany
4-3-3	Date of deposit	04 June 1999 (04.06.1999)
4-3-4	Accession Number	DSMZ 12844
4-4	Additional Indications	NONE
4-5	Designated States for Which Indications are Made	all designated States
4-6	Separate Furnishing of Indications	NONE
	These indications will be submitted to the International Bureau later	
5	The indications made below relate to	
	the deposited microorganism(s) or other biological material referred to in	
	the description on:	
5-1	page	5
5-2	line	7
5-3 5-3-1	Identification of Deposit Name of depositary institution	DSMZ-Deutsche Sammlung von
J-J-1	папьс от перозналу візицивоп	Mikroorganismen und Zellkulturen GmbH
5-3-2	Address of depositary institution	Mascheroder Weg 1b, D-38124
	-	Braunschweig, Germany
5-3-3	Date of deposit	04 June 1999 (04.06.1999)
5-3-4	Accession Number	DSMZ 12845
5-4	Additional Indications	NONE -
		ATVATAS

PCT

Original (for SUBMISSION) - printed on 30.08.2000 01:21:07 PM

5-5	Designated States for Which Indications are Made	all designated States
5-6	Separate Furnishing of Indications	NONE
	These indications will be submitted to the International Bureau later	
	FOR	RECEIVING OFFICE USE ONLY
0-4	This form was received with the international application: (yes or no)	RO/DK 3 1 AUG 2000
0-4-1	Authorized officer	Allui . Marie Louise Rosendal Head Clerk
	FOR INT	TERNATIONAL BUREAU USE ONLY
0-5	This form was received by the international Bureau on:	19.09.2000.
0-5-1	Authorized officer	13.03.2000.

Claims

	1. A RP-II protease selected from the group consisting of:
	(i) a RP-II protease that is immunochemically
5	identical or partially identical by cross-
	reaction with an antibody raised against or
	reactive with at least one epitope of a RP-I
	protease comprising the amino acid sequences
	of the mature peptides shown in the appended
10	Sequence Listing SEQ ID NO: 2, SEQ ID NO: 4
	SEQ ID NO: 6, SEQ ID NO: 8, SEQ ID NO: 10, or
	SEQ ID NO: 12; and/or
	(ii) a RP-II protease that is at least 60
	homologous with the amino acid sequence of a
15	RP-II protease comprising the amino acid
	sequence shown in the appended Sequence
	Listing SEQ ID NO: 2, SEQ ID NO: 4, SEQ ID
	NO: 6, SEQ ID NO: 8, SEQ ID NO: 10, or SEQ ID
	NO: 12; and/or
20	(iiia) a RP-II protease that is encoded by a DN
	sequence which hybridizes with a
	oligonucleotide probe hybridizing with a DN
	sequence encoding a RP-II protease comprising
	the amino acid sequence shown in the appended
25	Sequence Listing SEQ ID NO: 2, SEQ ID NO: 4
	SEQ ID NO: 6, SEQ ID NO: 8, SEQ ID NO: 10, or
	SEQ ID NO: 12; and/or
	(iiib) a RP-II protease that is encoded by a DNA sequence which hybridizes with a
2.0	sequence which hybridizes with a oligonucleotide probe hybridizing with a DN
30	sequence encoding a RP-II protease comprising
	the DNA sequence shown in the appended
	Sequence Listing SEQ ID NO: 1, SEQ ID NO: 3
	SEQ ID NO: 5, SEQ ID NO: 7, SEQ ID NO: 9, 0
35	SEQ ID NO: 11;
35	an allelic variant of (i), (ii), or (iiia or iiib);
	a subsequence of (i), (ii), (iiia or iiib), or (iv)
	wherein the subsequence has protease activity.

2. A RP-II protease variant, wherein at least one Asn and/or Gly in an Asn-Gly sequence has been modified by substitution, deletion and/or insertion to change or remove said Asn-Gly sequence.

5

- 3. The protease variant of claim 2, wherein the parent protease is selected from the group consisting of BLC, AC116, CDJ-31, BO32, JA96, AA513, MPR or variants thereof.
- 10 4. A RP-II protease variant, wherein any Glu and/or Asp residue has been modified by substitution or deletion.
 - 5. The protease variant of claim 4, wherein the parent protease is selected from the group consisting of BLC, AC116, CDJ-31, BO32, JA96, AA513, MPR or variants thereof.
 - 6. A RP-II protease variant wherein the amino acid residue occupying the 1st and/or 2nd position following a Glu or Asp has been modified by replacing said Glu or Asp residue with a Pro residue.
 - 7. The protease variant of claim 6, wherein the parent protease is selected from the group consisting of BLC, AC116, CDJ-31, BO32, JA96, AA513, MPR or variants thereof.

25

20

8. A RP-II protease variant wherein at least one amino acid residue susceptible to oxidation exposed to the surface of the molecule, is deleted or replaced with another amino acid residue less susceptible to oxidation.

30

- 9. The variant of claim 8, wherein the amino acid residue susceptible to oxidation is methionine.
- 10. The variant of claim 9 comprising a replacement by an acid residue selected from the group consisting of A, E, N, Q, I, L, S and K.
 - 11. The protease variant of any of claims 8, 9, or 10, wherein the parent protease is selected from the group

consisting of BLC, AC116, CDJ-31, BO32, JA96, AA513, MPR or variants thereof.

- 12. A RP-II protease variant wherein one or more tryptophan residues at the surface of the protease has been deleted or substituted.
 - 13. The variant of claim 12 comprising the modification of said tryptophan residues by substitution with F, T, Q or G.
- 14. The protease variant of any of claims 12, or 13, wherein the parent protease is selected from the group consisting of BLC, AC116, CDJ-31, BO32, JA96, AA513, MPR or variants thereof.
- 15. A RP-II protease variant wherein one or more tyrosine residues at the surface of the protease has been substituted.

15

25

- 16. The variant of claim 15 comprising the substitution of 20 said Tyr with Phe or Trp.
 - 17. The protease variant of any of claims 15, or 16, wherein the parent protease is selected from the group consisting of BLC, AC116, CDJ-31, BO32, JA96, AA513, MPR or variants thereof.
 - 18. A RP-II protease BLC variant comprising E152{R,K,G}.
- 19. A RP-II protease variant essentially as specifically30 disclosed herein.
 - 20. An isolated polynucleotide having a nucleic acid sequence encoding a RP-II protease, selected from the group consisting of:
- a nucleic acid sequence encoding a RP-II protease having an amino acid sequence which has an identity of at least 60% with an amino acid sequence of the mature polypeptide of SEQ ID NO: 2, 4, 6, 8, 10 or 12;

10

- (b) a nucleic acid sequence having at least 60% homology with a nucleic acid sequence encoding a mature RP-II protease of SEQ ID NO: 1, 3, 5, 7, 9, or 11;
- (c) a nucleic acid sequence which hybridizes
 under low stringency conditions with (i) the
 nucleic acid sequence of SEQ ID NO: 1, 3, 5,
 7, 9, or 11, (ii) the cDNA sequence of SEQ ID
 NO: 1, 3, 5, 7, 9, or 11, (iii) a subsequence
 of (i) or (ii) of at least 100 nucleotides,
 or (iv) a complementary strand of (i), (ii),
 or (iii);
- (d) an allelic variant of (a), (b), or (c);
 and/or
- 15 (e) a subsequence of (a), (b), (c), or (d), wherein the subsequence encodes a RP-II protease fragment which has protease activity.
- 20 21. The polynucleotide of claim 20, which encodes a RP-II protease having an amino acid sequence which has at least 60, 70, 80, 90, or 95% identity with the amino acid sequence of the mature peptide of SEQ ID NO: 2, 4, 6, 8, 10, 12, or 14, or corresponding subsequences thereof.
- 22. The polynucleotide of claim 20, which encodes a RP-II protease comprising the amino acid sequence of SEQ ID NO: 2, 4, 6, 8, 10, or 12.
- 30 23. The polynucleotide of claim 20, which encodes a RP-II protease consisting of the amino acid sequence of SEQ ID NO: 2, 4, 6, 8, 10 or 12, or a fragment thereof which has protease activity.
- 35 24. The polynucleotide of claim 28, which encodes a RP-II protease consisting of the amino acid sequence of the mature peptide of SEQ ID NO: 2, 4, 6, 8, 10 or 12.

- The polynucleotide of claim 20, which has at least 60, 70, 80, 90, or 95% homology with the nucleic acid sequence of SEQ ID NO: 1, 3, 5, 7, 9, or 11.
- 5 26. The polynucleotide of claim 20, which has the nucleic acid sequence of SEQ ID NO: 1, 3, 5, 7, 9, or 11.
- the claim 20, wherein 27. The polynucleotide of polynucleotide hybridizes under medium or high low, stringency conditions with (i) a polynucleotide comprising 10 the nucleic acid sequence of SEQ ID NO: 1, 3, 5, 7, 9, or 11, (ii) a polynucleotide comprising the cDNA sequence of SEQ ID NO: 1, 3, 5, 7, 9, or 11, (iii) a subsequence of (i) or (ii) of at least 100 nucleotides, or (iv) a complementary strand of (i), (ii), or (iii). 15
 - 28. The polynucleotide of claim 20, which is contained in the plasmid pUC19/AC116; pUC/CDJ31; pUC/BO32; pUC/JA96; or pUC/AA513 which is contained in DSM 12841: E. coli
- 20 pUC19/AC116; DSM 12842: E. coli pUC/CDJ31; DSM 12843: E. coli pUC/BO32; DSM 12844: E. coli pUC/JA96; or DSM 12845: E. coli pUC/AA513.
- 29. The polynucleotide of any of claims 20-28, which encodes a RP-II protease that has at least 20% of the protease activity of the RP-II protease of SEQ ID NO: 2, 4, 6, 8, 10, or 12.
- 30- An isolated polynucleotide comprising a nucleic acid sequence having at least one mutation in the mature RP-II protease coding sequence of SEQ ID NO: 1, 3, 5, 7, 9, 11, or 13, in which the mutant nucleic acid sequence encodes a mature polypeptide of SEQ ID NO: 2, 4, 6, 8, 10, 12 or 14.
- 31. An isolated polynucleotide produced by (a) hybridizing a DNA under low, medium or high stringency conditions with (i) the nucleic acid sequence of SEQ ID NO: 1, 3, 5, 7, 9, 11, or 13, (ii) the cDNA sequence of SEQ ID NO: 1, (iii) a subsequence of (i) or (ii) of at least 100 nucleotides, or

- (iv) a complementary strand or (i), (ii), or (iii); and (b) isolating the polynucleotide.
- 32. A nucleic acid construct comprising the polynucleotide of any of claims 20-31 operably linked to one or more control sequences that direct the production of the RP-II protease in a suitable expression host.
- 33. A recombinant expression vector comprising the nucleic acid construct of claim 32, a promoter, and transcriptional and translational stop signals.
 - 34. A recombinant host cell comprising the nucleic acid construct of claim 33.
- 35. A method for producing a mutant polynucleotide, comprising (a) introducing at least one mutation into the mature polypeptide coding sequence of SEQ ID NO: 1, 3, 5, 7, 9, 11, or 13, wherein the mutant polynucleotide encodes a RP-II protease consisting of SEQ ID NO: 2, 4, 6, 8, 10, 12 or 14; and (b) recovering the mutant polynucleotide.
 - 36. A mutant polynucleotide produced by the method of claim
 35.
 - 37. A method for producing a RP-II protease, comprising

 (i) cultivating a strain comprising the mutant polynucleotide of claim 36 encoding the RP-II protease to produce a supernatant comprising the RP-II protease; and
 - (ii) recovering the RP-II protease.

- - (ii) recovering the RP-II protease.
 - _ 39. A method for producing a RP-II protease comprising

- (i) cultivating a host cell under conditions conducive for production of the RP-II protease, wherein the host cell comprises the mutant polynucleotide of claim 36 encoding the RP-II protease to produce a supernatant comprising the polypeptide; and
- (ii) recovering the RP-II protease.
- 40. A nucleic acid construct comprising a gene encoding a protein operably linked to one or both of a first polynucleotide encoding a signal peptide from SEQ ID NO: 1, 3, 5, 7, 9, 11, or 13 and a second polynucleotide encoding a propeptide from SEQ ID NO: 1, 3, 5, 7, 9, 11, or 13, wherein the gene is foreign to the first and second polynucleotides.
- 15 41. A recombinant expression vector comprising the nucleic acid construct of claim 40.
 - 42. A recombinant host cell comprising the nucleic acid construct of claim 40 or the expression vector of claim 41.

5

- 43. A method for producing a protein comprising
 - (i) cultivating the recombinant host cell of claim 42 under conditions suitable for production of the protein; and
- 25 (ii) recovering the protein.

-, ---

- 44. A detergent composition comprising a RP-II protease of any of claims 1 to 19.
- 30 45. The composition of claim 44 comprising at least one further enzyme, preferably another protease, preferably a subtilisin of the subtilase group I-S2 or high alkaline subtilisins, preferably subtilisin PB92, subtilisin 309, subtilisin 147, or alkaline elastase YaB or functional variants thereof.
 - 46. The compositions of claim 44, wherein the other enzyme is a lipase, cellulase, amylase, peroxidase or oxidase.

- -

47. A method of using a RP-II protease of any of the claims 1 to 19 in a detergent composition.

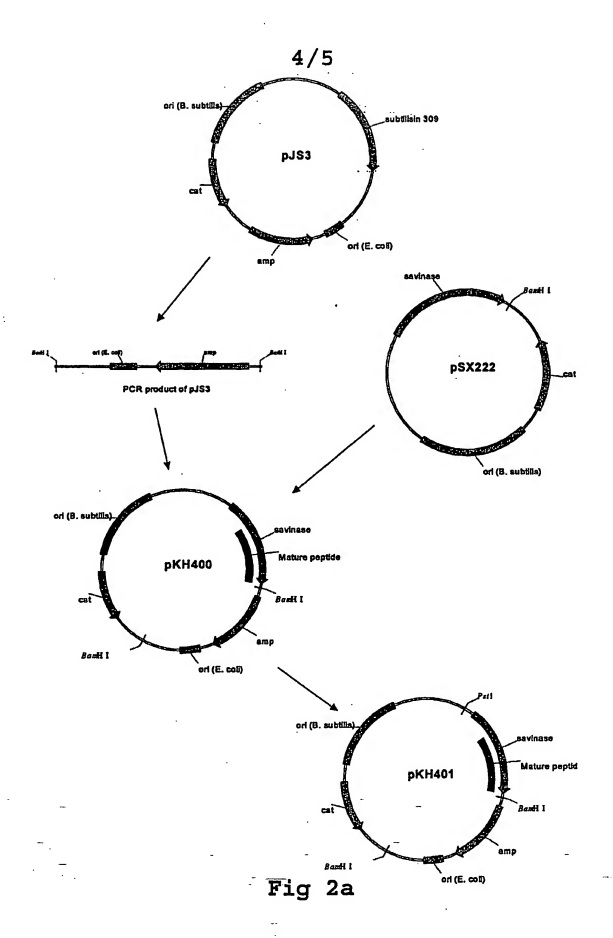
- a) BLC = RP-II protease from Bacillus licheniformis,
- b) AA513 = RP-II protease from Bacillus halmapalus AA513
- c) AC116 = RP-II protease from Bacillus licheniformis AC116
- d) BO32 = RP-II protease from Bacillus pumilus BO32
- e) CDJ31 = RP-II protease from Bacillus licheniformis CDJ31
- f) JA96 = RP-II protease from Bacillus pumilus JA96
- g) MPR = RP-II protease from Bacillus subtilis IS75
- a) 1 SVIGSDDRTRVTNTTAYPYRAIVHIS....SSIGSCTGWMIGPKTVATAG 46 VVIGDDGRRQVQNTSFMPFRALTYIEFGNLTSTWSCSGGVIGTDLVVTNA
- 47 HCIYDTSSGSFAGTATVSPGRNGTSYPYGSVKSTRYFIPSGWR.SGNTNY 95
- b) HCV....EGSVL.AGTVVPGMNNSQWAYGHYRVTQIIYPDQYRNNGASEF
- a) 96 DYGAIELS.....EPIGNTVGYFGYSYTTSSLVGTTVTISGYPGDKTAGT 140
- DYAILRVAPDSDGRHIGNRAGILSFTETGTVNENTFLRTYGYPGDKISET
- a) 141 Q....WQHSG.PIAISETYKLQYAMDTYGGQSGSPVFEQSSSRTNCSGPC 185
- KLISLWGMVGRSDAFLHRDLLFYNMDTYFGQSGSPVLNSVDSMVAVHNAG
- a) 186 SLAVHTNGVYGGSSYNRGTRITKEVFDNLTNWKNSAQ 222
- YIVGGNREINGGPKIRRDFTNLFNOMN......
- a) 1 SVIGSDDRTRVTNTTAYPYRAIVHISSSIGSCTGWMIGPKTVATAGHCIY 50
- SVIGSDERTRYTDTTAFPYRAIVHISSSIGSCTGWLIGPKTVATAGHCVY
- a) 51 DTSSGSFAGTATVSPGRNGTSYPYGSVKSTRYFIPSGWRSGNTNYDYGAI 100
- DTASRSFAGTATVSPGRNGSAYPYGSVTSTRYFIPSGWOSGNSNYDYAAI
- a) 101 ELSEPIGNTVGYFGYSYTTSSLVGTTVTISGYPGDKTAGTQWQHSGPIAI 150
- ELSQPIGNTVGYFGYSYTASSLAGAGVTISGYPGDKTTGTQWQMSGTIAV
- a) 151 SETYKLQYAMDTYGGQSGSPVFEQSSSRTNCSGPCSLAVHTNGVYGGSSY 200
- SETYKLQYAIDTYGGQSGSPVYEKSSSRTNCSGPCSLAVHTNGVYGGSSY c)
- a) 201 NRGTRITKEVFDNLTNWKNSAQ 222
- NRGTRITKEVFDNFTSWKNSAQ 222

Fig.

a) d)	1	SVIGSDDRTRVTNTTAYPYRAIVHISSSIGSCTGWMIGPKTVATAGHCIY VVIGDDGRTKVANTRVAPYNSIAYTTFGGSSCTGTLIAPNKILTNGHCVY	50
a) d)	51	DTSSGSFAGTATVSPGRNGTSYPYGSVKSTRYFIPSGW.RSGNTNYDYGA NTASRSYSAKGSVYPGMNDSTAVNGSANMTEFYVPSGYINTGASQYDFAV	99
a) d)	100	IELSEPIGNTVGYFGYSYTTSSLVGTTVTISGYPGDKTAGTQWQHSIKTDTNIGNTVGYRSIRQVT.NLTGTTIKISGYPGDKMRSTGKISQWEMS	145
a) d)	146	GPIAISETYKLQYAMDTYGGQSGSPVFEQSSSRTNC.SGPCSLAVHTNGV GPVTREDTNLAYYMIDTFSGNSGSAMLDQNQQIVGVHNAGYSNGTINGGP	194
a) d)	195	YGGSSYNRGTRITKEVFDNLTNWKNSAQ 222 KATAAFVEFINYAKAQ	
•	•		
a) e)	1	SVIGSDDRTRVTNTTAYPYRAIVHISSSIGSCTGWMIGPKTVATAGHCIY SVIGSDERTRVTNTTAYPYRAIVHISSSIGSCTGSLIGPKTVATAGHCIY	50
_ \			
a) e)	21	DTSSGSFAGTATVSPGRNGTSYPYGSVKSTRYFIPSGWRSGNTNYDYGAI DTASGSFAGTATVSPGRNGSTYPYGSVTSTRYFIPSGYRSGNSNYDYGAI	100
a)	101	ELSEPIGNTVGYFGYSYTTSSLVGTTVTISGYPGDKTAGTQWQHSGPIAI	150
e)		ELSQPIGNTVGYFGYSYTTSSLVGSSVTIIGYPGDKTSGTQWQMSGNIAV	
a)	151	SETYKLQYAMDTYGGQSGSPVFEQSSSRTNCSGPCSLAVHTNGVYGGSSY	200
e)		SETYKLQYAIDTYGGQSGSPVYEASSSRTNCSGPCSLAVHTNGVYGGSSY .	
a)	201	NRGTRITKEVFDNLTNWKNSAQ 222	
e)		NRGTRITKEVFDNLTNWKNSAQ	
	•		
a)	1	SVIGSDDRTRVTNTTAYPYRAIVHISSSIGSCTGWMIGPKTVATAGHCIY	50
f)	_	VVIGDDGRTKVTNTRVAPYNSIAYITFGGSSCTGTLIAPNKILTNGHCVY	
a)	51	DTSSGSFAGTATVSPGRNGTSYPYGSVKSTRYFIPSGW.RSGNTNYDYGA	99
f)		NTATRSYSAKGSVYPGMNDSTAVNGSANMTEFYVPSGYINTGASQYDFAV	
a) f)	100	IELSEPIGNTVGYFGYSYTTSSLVGTTVTISGYPGDKTAGTQWQHS IKTDTNIGNTVGYRSIRQVT.NLTGTTIKISGYPGDKMRSTGKVSQWEMS	145
a) f)	146	GPIAISETYKLQYAMDTYGGQSGSPVFEQSSSRTNC.SGPCSLAVHTNGV GPVTREDTNLAYYTIDTFSGNSGSAMLDQNQQIVGVHNAGYSNGTINGGP	194
a)	195	YGGSSYNRGTRITKEVFDNLTNWKNSAQ 222	
E)		KATAAFVEFINYAKAQ	

93
142
192

WO 01/16285 PCT/DK00/00476.



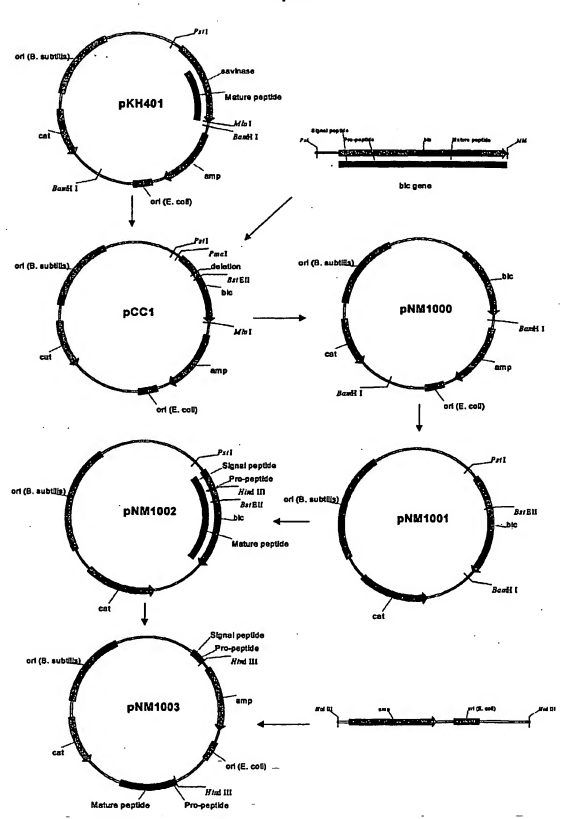


Fig 2b

SEQUENCE LISTING

<110> Novo Nordisk A/S

<120> Novel Proteases And Variants Thereof

<130> 5665.204-WO

<140>
<141>

<150> PA 1999 01212
<151> 1999-08-31

<150> PA 1999 01500
<151> 1999-10-20

<160> 45

<170> PatentIn Ver. 2.1

<210> 1

\Z1U> 1

<211> 948

<212> DNA

<213> Bacillus licheniformis

<220>

<221> CDS

<222> (1)..(948)

<220>

<221> mat_peptide

<222> (283)..(948)

<220>

<221> sig_peptide

<222> (1)..(93)

<223> pro_peptide (94) ... (282)

<400> 1

-3-

ttg gtt agt aaa aag agt gtt aaa cga ggt ttg atc aca ggt ctc att 48
Leu Val Ser Lys Lys Ser Val Lys Arg Gly Leu Ile Thr Gly Leu Ile
-90 -85 -80

ggt att tet att tat tet tta ggt atg cac eeg gee caa gee geg eea 96

Gly	Ile	Ser	Ile -75	Tyr	Ser	Leu	Gly	Met -70	His	Pro	Ala	Gln	Ala -65	Ala	Pro	
_					_		-	_						gaa Glu		144
_	-			_			_	-	_	-	_			ttg Leu		192
					_		_					_		gaa Glu		240
														tcg Ser 1		288
			_	-				_					-	tat Tyr	-	336
	-			•				-	-				_	acc Thr		384
	•			_			_	-		-			_	atc Ile		432
_			_				_			-		_	_	ccg Pro 65		480
				_						_		_	_	cgc Arg		528
		_				_	_						_	tac Tyr		576
														ttc Phe		624
tac	tcg	tac	act	act	tca	tca	ctt	gtt	ggg	aca	act	gtt	acc	atc	agc	672

Tyr Ser Tyr Thr Thr Ser Ser Leu Val Gly Thr Thr Val Thr Ile Ser 120 125 115 ggc tac cca ggc gat aaa aca gca ggc aca caa tgg cag cat tca gga Gly Tyr Pro Gly Asp Lys Thr Ala Gly Thr Gln Trp Gln His Ser Gly 145 135 768 ccg att gcc atc tcc gaa acg tat aaa ttg cag tac gca atg gac acg Pro Ile Ala Ile Ser Glu Thr Tyr Lys Leu Gln Tyr Ala Met Asp Thr 150 155 160 816 tac gga gga caa agc ggt tca ccg gta ttc gaa caa agc agc tcc aga Tyr Gly Gly Gln Ser Gly Ser Pro Val Phe Glu Gln Ser Ser Arg 170 175 165 864 acg aac tgt agc ggt ccg tgc tcg ctt gcc gta cac aca aat gga gta Thr Asn Cys Ser Gly Pro Cys Ser Leu Ala Val His Thr Asn Gly Val 180 185 190 tac ggc ggc tcc tcg tac aac aga ggc acc cgg att aca aaa gag gtg Tyr Gly Gly Ser Ser Tyr Asn Arg Gly Thr Arg Ile Thr Lys Glu Val 205 210 195 200 948 ttc gac aat ttg acc aac tgg aaa aac agc gca caa Phe Asp Asn Leu Thr Asn Trp Lys Asn Ser Ala Gln 215 220 <210> 2 <211> 316 <212> PRT <213> Bacillus licheniformis Leu Val Ser Lys Lys Ser Val Lys Arg Gly Leu Ile Thr Gly Leu Ile -85 -80 -90 Gly Ile Ser Ile Tyr Ser Leu Gly Met His Pro Ala Gln Ala Ala Pro -75 -70 -65 Ser Pro His Thr Pro Val Ser Ser Asp Pro Ser Tyr Lys Ala Glu Thr -60 -55 -50 Ser Val Thr Tyr Asp Pro His Ile Lys Ser Asp Gln Tyr Gly Leu Tyr -40 -35 -45 Ser Lys Ala Phe Thr Gly Thr Gly Lys Val Asn Glu Thr Lys Glu Lys -20 -25

Ala Glu Lys Lys Ser Pro Ala Lys Ala Pro Tyr Ser Ile Lys Ser Val -10 -5 Ile Gly Ser Asp Asp Arg Thr Arg Val Thr Asn Thr Thr Ala Tyr Pro Tyr Arg Ala Ile Val His Ile Ser Ser Ser Ile Gly Ser Cys Thr Gly Trp Met Ile Gly Pro Lys Thr Val Ala Thr Ala Gly His Cys Ile Tyr Asp Thr Ser Ser Gly Ser Phe Ala Gly Thr Ala Thr Val Ser Pro Gly Arg Asn Gly Thr Ser Tyr Pro Tyr Gly Ser Val Lys Ser Thr Arg Tyr Phe Ile Pro Ser Gly Trp Arg Ser Gly Asn Thr Asn Tyr Asp Tyr Gly Ala Ile Glu Leu Ser Glu Pro Ile Gly Asn Thr Val Gly Tyr Phe Gly Tyr Ser Tyr Thr Thr Ser Ser Leu Val Gly Thr Thr Val Thr Ile Ser Gly Tyr Pro Gly Asp Lys Thr Ala Gly Thr Gln Trp Gln His Ser Gly Pro Ile Ala Ile Ser Glu Thr Tyr Lys Leu Gln Tyr Ala Met Asp Thr Tyr Gly Gly Gln Ser Gly Ser Pro Val Phe Glu Gln Ser Ser Ser Arg Thr Asn Cys Ser Gly Pro Cys Ser Leu Ala Val His Thr Asn Gly Val · 185 Tyr Gly Gly Ser Ser Tyr Asn Arg Gly Thr Arg Ile Thr Lys Glu Val Phe Asp Asn Leu Thr Asn Trp Lys Asn Ser Ala Gln ⁻ 215 ₋

```
<210> 3
<211> 1026
<212> DNA
<213> Bacillus halmapalus AA513
<220>
<221> CDS
<222> (1)..(1026)
<220>
<221> mat_peptide
<222> (361)..(1026)
<220>
<221> sig_peptide
<222> (1)..(78)
<223> Pro-peptide (79) ... (360)
<400> 3
atg aaa cta cta tta aaa ctt act ttt gta tgc ata ttt atg tta agt
Met Lys Leu Leu Lys Leu Thr Phe Val Cys Ile Phe Met Leu Ser
                   -115
-120
                                       -110
ggg att cta tcc cca gta aac gca act caa gct gag act ctt act aaa
                                                                   96
Gly Ile Leu Ser Pro Val Asn Ala Thr Gln Ala Glu Thr Leu Thr Lys
               -100
                                    -95
tta aat aaa ata agt cag aag cag gaa cca tca tat aaa cta gat gaa
                                                                   144
Leu Asn Lys Ile Ser Gln Lys Gln Glu Pro Ser Tyr Lys Leu Asp Glu
            -85
                                -80
                                                                   192
gaa atg gat tat gtt cta att gat ttg gaa aca caa tct gaa tcg att
Glu Met Asp Tyr Val Leu Ile Asp Leu Glu Thr Gln Ser Glu Ser Ile
        -70
                            -65
                                                -60
att tcg ata gga gat aat acc gat ttg gga gat caa tcg ttt act tct
                                                                   240
Ile Ser Ile Gly Asp Asn Thr Asp Leu Gly Asp Gln Ser Phe Thr Ser
    -55
                        -50
tta ggg aag gtg gga cat gga gaa ctt gag aaa att aac tta gaa gaa
Leu Gly Lys Val Gly His Gly Glu Leu Glu Lys Ile Asn Leu Glu Glu
-40
                    -35
                                        -30
ttt cgt aat cct aat tta aca gta gta gac ccg tta aca cgt aag cct
                                                                   336
Phe Arg Asn Pro Asn Leu Thr Val Val Asp Pro Leu Thr Arg Lys Pro
                                    -15
                -20
att gaa caa aaa atc agc cct ttt gtt gtt ata ggc gat gat ggg aga
                                                                   384
```

_

Ile	Glu	Gln	Lys -5	Ile	Ser	Pro	Phe -1	Val 1	Val	Ile	Gly	Asp 5	Asp	Gly	Arg	
		gtt Val														432
	-	ttt Phe														480
		aca Thr														528
		gca Ala														576
		cat His 75														624
		ggt Gly														672
		gat Asp														720
	-	aca Thr														768
		ggt Gly														816
_	-	ggt Gly 155	_													864
		gac Asp														912
			a to c-		~~~	~++	ast	22+	~~~	~~~	tat	atc	att	aat	aat	960

Val Asp Ser Met Val Ala Val His Asn Ala Gly Tyr Ile Val Gly 185 190 195 aat agg gaa att aat ggt ggt cct aaa atc aga aga gat ttt aca aac Asn Arg Glu Ile Asn Gly Gly Pro Lys Ile Arg Arg Asp Phe Thr Asn 205 210 1026 tta ttt aat caa atg aac Leu Phe Asn Gln Met Asn 220 <210> 4 <211> 342 <212> PRT <213> Bacillus halmapalus AA513 <400> 4 Met Lys Leu Leu Lys Leu Thr Phe Val Cys Ile Phe Met Leu Ser -120 -115 -110 -105 Gly Ile Leu Ser Pro Val Asn Ala Thr Gln Ala Glu Thr Leu Thr Lys -100 -95 Leu Asn Lys Ile Ser Gln Lys Gln Glu Pro Ser Tyr Lys Leu Asp Glu -85 -80 Glu Met Asp Tyr Val Leu Ile Asp Leu Glu Thr Gln Ser Glu Ser Ile -70 -65 -60 Ile Ser Ile Gly Asp Asn Thr Asp Leu Gly Asp Gln Ser Phe Thr Ser -50 Leu Gly Lys Val Gly His Gly Glu Leu Glu Lys Ile Asn Leu Glu Glu -35 -30 Phe Arg Asn Pro Asn Leu Thr Val Val Asp Pro Leu Thr Arg Lys Pro -20 -15 Ile Glu Gln Lys Ile Ser Pro Phe Val Val Ile Gly Asp Asp Gly Arg -5 -1 1 Arg Gln Val Gln Asn Thr Ser Phe Met Pro Phe Arg Ala Leu Thr Tyr 10 15 20 Ile Glu Phe Gly Asn Leu Thr Ser Thr Trp Ser Cys Ser Gly Gly Val _ 30 35

Ile Gly Thr Asp Leu Val Val Thr Asn Ala His Cys Val Glu Gly Ser
45 50 55

Val Leu Ala Gly Thr Val Val Pro Gly Met Asn Asn Ser Gln Trp Ala 60 65 70

Tyr Gly His Tyr Arg Val Thr Gln Ile Ile Tyr Pro Asp Gln Tyr Arg
75 80 85

Asn Asn Gly Ala Ser Glu Phe. Asp Tyr Ala Ile Leu Arg Val Ala Pro 90 95 100

Asp Ser Asp Gly Arg His Ile Gly Asn Arg Ala Gly Ile Leu Ser Phe 105 110 115 120

Thr Glu Thr Gly Thr Val Asn Glu Asn Thr Phe Leu Arg Thr Tyr Gly 125 130 135

Tyr Pro Gly Asp Lys Ile Ser Glu Thr Lys Leu Ile Ser Leu Trp Gly
140 145 150

Met Val Gly Arg Ser Asp Ala Phe Leu His Arg Asp Leu Leu Phe Tyr 155 160 165

Asn Met Asp Thr Tyr Phe Gly Gln Ser Gly Ser Pro Val Leu Asn Ser 170 175 180

Val Asp Ser Met Val Ala Val His Asm Ala Gly Tyr Ile Val Gly Gly 185 190 195 200

Asn Arg Glu Ile Asn Gly Gly Pro Lys Ile Arg Arg Asp Phe Thr Asn 205 210 215

Leu Phe Asn Gln Met Asn 220

<210> 5

<211> 942

<212> DNA

<213> Bacillus licheniformis AC116

<220>

<221> CDS

<222> (1)..(942)

<220>

<221> mat_peptide <222> (277)..(942) <220> <221> sig_peptide <222> (1)..(87) <223> pro-peptide (88) ... (276) <400> 5 atg gcg aaa aat ggt gtt tca cgc gtt ttc att gcc gga ctc atc gga Met Ala Lys Asn Gly Val Ser Arg Val Phe Ile Ala Gly Leu Ile Gly -90 -80 -85 att tet att ttt tet teg gge att tac tet gea caa get gea tea teg 96 Ile Ser Ile Phe Ser Ser Gly Ile Tyr Ser Ala Gln Ala Ala Ser Ser -70 ceg cat acc cca gtc tcc agc gac cct tcg tac aag ccc ggc tcc acc Pro His Thr Pro Val Ser Ser Asp Pro Ser Tyr Lys Pro Gly Ser Thr -60 -55 -50 -45 tat gat ccc aac ata aaa att gac aat aac ggc gca tat tcg aaa gcc 192 Tyr Asp Pro Asn Ile Lys Ile Asp Asn Asn Gly Ala Tyr Ser Lys Ala -40 -35 ttc gaa gga acc gga aca ccc ggc ggc tcc gtt cag gcc aaa ccg aaa 240 Phe Glu Gly Thr Gly Thr Pro Gly Gly Ser Val Gln Ala Lys Pro Lys -25 -20 aaa gaa tcg ccc gcc ggc ccg cct tac agc cct aaa tcg gta atc ggc 288 Lys Glu Ser Pro Ala Gly Pro Pro Tyr Ser Pro Lys Ser Val Ile Gly -5 -10 tca gat gaa cgg aca agg gtg act gat aca acg gcc ttt cca tac aga Ser Asp Glu Arg Thr Arg Val Thr Asp Thr Thr Ala Phe Pro Tyr Arg 5 10 20 gca atc gtc cat att tca agc agc atc ggc tca tgc aca ggc tgg ctg 384 Ala Ile Val His Ile Ser Ser Ser Ile Gly Ser Cys Thr Gly Trp Leu 25 30 432 atc gga ccg aaa acg gta gca acg gcc ggg cac tgc gtc tat gac acg Ile Gly Pro Lys Thr Val Ala Thr Ala Gly His Cys Val Tyr Asp Thr 40 45 50 gca agc cga tca ttc gcg gga acc gcc acc gtt tcc ccg gga cga aac 480 Ala Ser Arg Ser Phe Ala Gly Thr Ala Thr Val Ser Pro Gly Arg Asn 55 60 65

PCT/DK00/00476 WO 01/16285

ggt	tca	gct	tac	cct	tac	gga	tct	gtt	aca	tcg	acc	cgc	tat	ttc	atc	528
Gly	Ser	Ala	Tyr	Pro	Tyr	Gly	Ser	Val	Thr	Ser	Thr	Arg	Tyr	Phe	Ile	
_	70					75					80					
cca	tca	aat	taa	cag	agc	qqa	aat	tcc	aat	tat	gac	tac	gca	gcg	atc	576
				Gln												
85		,		•=	90	2		-		95	•	•			100	
•					•											
gag	ctc	agc	cag	ccg	atc	ggc	aat	acc	atc	σσа	tat	ttc	aga	tat	tca	624
		-	_	Pro												
Giu	Deu	Der	J 111	105		0.7			110	0-3	-3-		1	115		
				103												
tac	200	act	tca	tcg	ctt	aca.	aaa	σca	aac	ata	acc	atc	age	gga	tat	672
		_		Ser		_		-								• • •
ıyı	1111	AIG		Ser	пец	AIG	GIJ	125	GLY	Val	1111	110	130	013	-3-	
			120					123					130			
													~~-	200	250	720
				aca												120
Pro	GīĀ		rys	Thr	THE	GIY		GIN	тр	GIN	met	_	GIA	THE	TTE	
		135					140					145				
																7.00
-	-			acg												768
Ala	Val	Ser	Glu	Thr	Tyr	Lys	Leu	Gln	Tyr	Ala		Asp	Thr	Tyr	Gly	
	150					155					160					
		-		tcc												816
Gly	Gln	Ser	Gly	Ser	Pro	Val	Tyr	Glu	Lys	Ser	Ser	Ser	Arg	Thr	Asn	
165					170					175					180	
tgc	agc	ggc	cca	tgc	tcg	ctg	gcc	gtt	cat	acg	aac	ggc	gtg	tac	gga	864
Cys	Ser	Gly	Pro	Cys	Ser	Leu	Ala	Val	His	Thr	Asn	Gly	Val	Tyr	Gly	
				185					190					195		
gga	tcc	tct	tac	aac	aga	ggc	acc	cgc	att	acg	aaa	gaa	gta	ttt	gat	912
Gly	Ser	Ser	Tyr	Asn	Arg	Gly	Thr	Arg	Ile	Thr	Lys	Glu	Val	Phe	Asp	
			200					205					210			
aat	ttc	aca	agc	tgg	aaa	aac	agc	gca	cag							942
			_	Trp												
		215		_			220									
<210)> 6															

<210> 6

<211> 314 <212> PRT ---

<213> Bacillus licheniformis AC116

<400> 6 --* ***

Met Ala Lys Asn Gly Val Ser Arg Val Phe Ile Ala Gly Leu Ile Gly
-90 -85 -80

- Ile Ser Ile Phe Ser Ser Gly Ile Tyr Ser Ala Gln Ala Ala Ser Ser
 -75 -70 -65
- Pro His Thr Pro Val Ser Ser Asp Pro Ser Tyr Lys Pro Gly Ser Thr
 -60 -55 -50 -45
- Tyr Asp Pro Asn Ile Lys Ile Asp Asn Asn Gly Ala Tyr Ser Lys Ala
 -40 -35 -30
- Phe Glu Gly Thr Gly Thr Pro Gly Gly Ser Val Gln Ala Lys Pro Lys
 -25
 -20
 -15
- Lys Glu Ser Pro Ala Gly Pro Pro Tyr Ser Pro Lys Ser Val Ile Gly
 -10 -5 -1 1
- Ser Asp Glu Arg Thr Arg Val Thr Asp Thr Thr Ala Phe Pro Tyr Arg
 5 10 15 20
- Ala Ile Val His Ile Ser Ser Ser Ile Gly Ser Cys Thr Gly Trp Leu 25 30 35
- Ile Gly Pro Lys Thr Val Ala Thr Ala Gly His Cys Val Tyr Asp Thr
 40 45 50
- Ala Ser Arg Ser Phe Ala Gly Thr Ala Thr Val Ser Pro Gly Arg Asn
 55 60 65
- Gly Ser Ala Tyr Pro Tyr Gly Ser Val Thr Ser Thr Arg Tyr Phe Ile
 70 75 80
- Pro Ser Gly Trp Gln Ser Gly Asn Ser Asn Tyr Asp Tyr Ala Ala Ile 85 90 95 100
- Glu Leu Ser Gln Pro Ile Gly Asn Thr Val Gly Tyr Phe Gly Tyr Ser 105 110 115
- Tyr Thr Ala Ser Ser Leu Ala Gly Ala Gly Val Thr Ile Ser Gly Tyr 120 125 130
- Pro Gly Asp Lys Thr Thr Gly Thr Gln Trp Gln Met Ser Gly Thr Ile 135 140 145
- Ala Val Ser Glu Thr Tyr Lys Leu Gln Tyr Ala Ile Asp Thr Tyr Gly
 -150 155 160 ---

Gly Gln Ser Gly Ser Pro Val Tyr Glu Lys Ser Ser Ser Arg Thr Asn

165 . 170 175 Cys Ser Gly Pro Cys Ser Leu Ala Val His Thr Asn Gly Val Tyr Gly 190 185 Gly Ser Ser Tyr Asn Arg Gly Thr Arg Ile Thr Lys Glu Val Phe Asp 200 205 Asn Phe Thr Ser Trp Lys Asn Ser Ala Gln 215 220 <210> 7 <211> 909 <212> DNA <213> Bacillus pumilus BO32 <220> <221> CDS <222> (1)..(909) <220> <221> mat_peptide <222> (265)..(909) <220> <221> sig_peptide <222> (1)..(78) <223> pro-peptide (79) ... (264) <400> 7 atg atg aaa aag gtg aaa atg tta ctc cct tct cta ctt gtt ttt ggt Met Met Lys Lys Val Lys Met Leu Leu Pro Ser Leu Leu Val Phe Gly -75 -85 -80 gct tta agt gtg cct agt ttt gcc cat gcc gca tct gat tca gtg cta 96 Ala Leu Ser Val Pro Ser Phe Ala His Ala Ala Ser Asp Ser Val Leu -70 -65 acg tct gat tat gac atg gtg act tct gat gga aag gtg atc tct tca 144 Thr Ser Asp Tyr Asp Met Val Thr Ser Asp Gly Lys Val Ile Ser Ser -55 -50 agt gat ttc cac aat gat acg aaa tcc ccc tca tcc ttt gat aaa gtg 192 Ser Asp Phe His Asn Asp Thr Lys Ser Pro Ser Ser Phe Asp Lys Val **-35** --25 -30

12

		cta														240
qaA	qaA	Leu	Ser		Thr	Val	Gly	Glu		Val	Lys	Pro	Leu		Lys	
				-20					-15					-10		
															200	288
		aaa	_													200
ıyr	Leu	Lys		Pne	Gin	inr	Lys -1	val	vaı	TIE	GTÅ	ASP 5	Asp	GTÅ	Arg	
			-5				-1					,				
202	222	gta	aca	aat	aca	ana	ata	gca	cca	tat	aat	tca	att	act	tat	336
		Val														
	10					15					20				-	
act	acg	ttt	ggc	ggc	tcc	agc	tgc	acg	ggg	acc	ctg	att	gcc	cct	aac	384
Thr	Thr	Phe	Gly	Gly	Ser	Ser	Cys	Thr	Gly	Thr	Leu	Ile	Ala	Pro	Asn	
25					30					35					40	
		ttg														432
ГЛЗ	Ile	Leu	Thr		Gly	His	Cys	Val		Asn	Thr	Ala	Ser		Ser	
				45					50					55		
													-a-	201	~~~	480
		gca Ala														400
Tyr	ser	Ala	ьув 60	GIĀ	ser	vai	TYL	65	GIY	Mec	ASII	Asp	70	1111	ΛIα	
			00					0,5					, ,			
ata	aat	ggc	tca	gca	aat	atg	aca	gag	ttc	tat	gta	cca	agc	ggg	tat	528
		Gly														
		75					80					85				
atc	aat	aca	ggt	gcg	agc	çaa	tat	gat	ttt	gcc	gtg	atc	aaa	aca	gat	576
Ile	Asn	Thr	Gly	Ala	Ser	Gln	Tyr	Asp	Phe	Ala	Val	Ile	ГЛа	Thr	Asp	
	90					95					100					
		att												_	_	624
	Asn	Ile	GIA	Asn		vaı	GIA	туr	Arg		Tie	Arg	GIN	vaı	120	
105					110					115					120	
220	++=	act	aaa	909	aca	att	222	att	tct	gga	tat	cca	aat.	gat	aaa	672
		Thr											_			
71511	200		G _J	125			_, _		130	3	-4 -			135	•	
atg	aga	tca	act	ggc	aag	atc	tcg	cag	tgg	gag	atg	tça	ggt	cct	gtg	720
		Ser														
			140					145					150			
						-										_
	_	gaa														768
Thr	Arg	Glu	Asp	Thr	Asn	Leu		Tyr	Tyr	Met	Ile		Thr	Phe	Ser	
		155					160					165				

gga aat tca ggc tca gcg atg cta gat caa aat cag caa att gtt ggg Gly Asn Ser Gly Ser Ala Met Leu Asp Gln Asn Gln Gln Ile Val Gly 175 170 gtt cat aac gca ggg tat tca aac ggt acg att aat ggc ggt cca aaa Val His Asn Ala Gly Tyr Ser Asn Gly Thr Ile Asn Gly Gly Pro Lys 185 190 909 gcg aca gct gcc ttt gtt gaa ttt atc aac tat gca aaa gcg caa Ala Thr Ala Ala Phe Val Glu Phe Ile Asn Tyr Ala Lys Ala Gln 205 210 <210> 8 <211> 303 <212> PRT <213> Bacillus pumilus BO32 <400> 8 Met Met Lys Lys Val Lys Met Leu Leu Pro Ser Leu Leu Val Phe Gly -85 -80 Ala Leu Ser Val Pro Ser Phe Ala His Ala Ala Ser Asp Ser Val Leu -70 -65 Thr Ser Asp Tyr Asp Met Val Thr Ser Asp Gly Lys Val Ile Ser Ser -55 -50 Ser Asp Phe His Asn Asp Thr Lys Ser Pro Ser Ser Phe Asp Lys Val -40 -35 -30 Asp Asp Leu Ser Ser Thr Val Gly Glu Lys Val Lys Pro Leu Ser Lys -20 -15 Tyr Leu Lys Asp Phe Gln Thr Lys Val Val Ile Gly Asp Asp Gly Arg -5 -1 1 Thr Lys Val Ala Asn Thr Arg Val Ala Pro Tyr Asn Ser Ile Ala Tyr 20 10 15 Thr Thr Phe Gly Gly Ser Ser Cys Thr Gly Thr Leu Ile Ala Pro Asn 25 30 Lys Ile Leu Thr Asn Gly His Cys Val Tyr Asn Thr Ala Ser Arg Ser 50 Tyr Ser Ala Lys Gly Ser Val Tyr Pro Gly Met Asn Asp Ser Thr Ala

60 . 65 70

Val Asn Gly Ser Ala Asn Met Thr Glu Phe Tyr Val Pro Ser Gly Tyr
75 80 85

Ile Asn Thr Gly Ala Ser Gln Tyr Asp Phe Ala Val Ile Lys Thr Asp 90 95 100

Thr Asn Ile Gly Asn Thr Val Gly Tyr Arg Ser Ile Arg Gln Val Thr 105 110 115 120

Asn Leu Thr Gly Thr Thr Ile Lys Ile Ser Gly Tyr Pro Gly Asp Lys 125 130 135

Met Arg Ser Thr Gly Lys Ile Ser Gln Trp Glu Met Ser Gly Pro Val 140 145 150

Thr Arg Glu Asp Thr Asn Leu Ala Tyr Tyr Met Ile Asp Thr Phe Ser 155 160 165

Gly Asn Ser Gly Ser Ala Met Leu Asp Gln Asn Gln Gln Ile Val Gly 170 175 180

Val His Asn Ala Gly Tyr Ser Asn Gly Thr Ile Asn Gly Gly Pro Lys
185 190 195 200

Ala Thr Ala Ala Phe Val Glu Phe Ile Asn Tyr Ala Lys Ala Gln
205 210 215

<210> 9

<211> 954

<212> DNA

<213> Bacillus licheniformis CDJ31

<220>

<221> CDS

<222> (1)..(954)

<220>

<221> mat_peptide

<222> (289)..(954)

<220>

<221> sig_peptide

<222> (1)..(84)

<223> pro-peptide (85) ... (288)

- ==-

	<400)> 9																
	atg	aaa	aaa	agt	gtg	aca	cgc	gta	tta	atg	gcc	ggt	ctt	att	gga	ata	48	
	Met	Lys	Lys	Ser	Val	Thr	Arg	Val	Leu	Met	Ala	Gly	Leu	Ile	Gly	Ile		
		-95					-90					-85						
	tct	att	tat	tct	atg	ggc	atc	gac	tcc	gct	caa	gct	gca	tca	tcg	ccg	96	
	Ser	Ile	Tyr	Ser	Met	Gly	Ile	Asp	Ser	Ala	Gln	Ala	Ala	Ser	Ser	Pro		
	-80					-75					-70					-65		
	cat	act	cct	gtc	tct	agc	gat	cct	tca	tac	aag	ccc	gac	tca	tcc	gca	144	
	His	Thr	Pro	Val	Ser	Ser	Asp	Pro	Ser	Tyr	Lys	Pro	Asp	Ser	Ser	Ala		
					-60					-55					-50			
	agc	tat	gat	cct	gct	att	aaa	acc	aac	aaa	aac	ggc	gcc	tat	tca	aaa	192	
	Ser	Tyr	Asp	Pro	Ala	Ile	Lys	Thr	Asn	Lys	Asn	Gly	Ala	Tyr	Ser	Lys		
				-45					-40					-35				
	gca	ttt	gaa	ggt	aca	gga	aaa	cta	gaç	gct	ccc	ctt	tat	cag	gaa	aaa	240	
	Ala	Phe	Glu	Gly	Thr	Gly	Lys	Leu	Asp	Ala	Pro	Leu	Tyr	Gln	Glu	Lys		
			-30					-25					-20					
	agc	aaa	cca	acc	aaa	aaa	tcc	cct	gcc	gga	cca	cgt	tac	agc	ccc	aaa	288	
						Lys												
		-15			_	_	-10			_		-5				-1		
	tcc	gtg	att	ggt	tct	gat	gaa	cgg	acg	aga	gtg	aca	aac	act	acc	gca	336	
	Ser	Val	Ile	Gly	Ser	Asp	Glu	Arg	Thr	Arg	Val	Thr	Asn	Thr	Thr	Ala		
	1			-	5	_		_		10					15			
	tat	cca	tac	aga	gcg	atc	gtg	cat	att	tca	agc	agc	atc	ggg	tct	tgc	384	
						Ile												
	•		•	20					25					30		-		
	acc	aac	tcc	cta	atc	ggt	ccg	aaa	acg	gtg	gca	acg	gcc	gga	cac	tgc	432	
						Gly												
		-	35			_		40					45					
	att	tat	gac	aca	aca	agc	aaa	tca	ttc	gcc	gga	acc	gct	acc	gtt	tct	480	
						Ser												
		50					55				_	60						
	cca	σσa	caa	aac	ggt	tca	aca	tat	ccg	tac	gga	tca	gtt	aca	tca	acc	528	
	_	_				Ser												
	65	1			-	70	_	•		•	75					80		
	cac	tat	ttc	atc	cca	tca	ggc	tat	cga	agc	gga	aat	tcg	aat	tac	gac	576	
-	_					Ser												
	>	- 2 -						_	- 0	_	-				_	_		

85 90 95

tac gga gcc ata gag ctc agc cag ccg atc ggc aac acc gtc ggg tat Tyr Gly Ala Ile Glu Leu Ser Gln Pro Ile Gly Asn Thr Val Gly Tyr 105 100 ttc gga tat tcc tac acc tcg tct ctc gtt ggg tca agc gtt acc Phe Gly Tyr Ser Tyr Thr Thr Ser Ser Leu Val Gly Ser Ser Val Thr 115 120 atc atc gga tat cca ggc gac aaa aca tcg ggc acc caa tgg cag atg Ile Ile Gly Tyr Pro Gly Asp Lys Thr Ser Gly Thr Gln Trp Gln Met 135 130 tcc gga aat atc gcc gtc tca gaa aca tat aaa ctg caa tat gcg atc Ser Gly Asn Ile Ala Val Ser Glu Thr Tyr Lys Leu Gln Tyr Ala Ile 150 145 gac aca tac gga ggg cag agc ggc tct ccc gta tat gag gcg agc agc Asp Thr Tyr Gly Gly Gln Ser Gly Ser Pro Val Tyr Glu Ala Ser Ser 165 170 tcc aga acg aat tgc agc ggc cca tgt tcg ctg gcc gtt cat acg aat Ser Arg Thr Asn Cys Ser Gly Pro Cys Ser Leu Ala Val His Thr Asn 185 180 ggg gtg tac gga gga tct tca tac aac aga ggc acc cgg att aca aaa Gly Val Tyr Gly Gly Ser Ser Tyr Asn Arg Gly Thr Arg Ile Thr Lys 195 200 205

gaa gta ttc gat aat ttg aca aac tgg aaa aac agc gcc caa 954
Glu Val Phe Asp Asn Leu Thr Asn Trp Lys Asn Ser Ala Gln
210 215 220

<210> 10

<211> 318

<212> PRT

<213> Bacillus licheniformis CDJ31

<400> 10

Met Lys Lys Ser Val Thr Arg Val Leu Met Ala Gly Leu Ile Gly Ile
-95 -90 -85

Ser Ile Tyr Ser Met Gly Ile Asp Ser Ala Gln Ala Ala Ser Ser Pro
-80 -75 -70 -65

His Thr Pro Val Ser Ser Asp Pro Ser Tyr Lys Pro Asp Ser Ser Ala

				-60					-55					-50	
Ser	Tyr	Asp	Pro -45	Ala	Ile	Lys	Thr	Asn -40	Lys	Asn	Gly	Ala	Tyr -35	Ser	Lys
Ala	Phe	Glu -30	Gly	Thr	Gly	Lys	Leu -25	Asp	Ala	Pro	Leu	Tyr -20	Gln	Glu	Lys
Ser	Lys -15	Pro	Thr	Lys	Lys	Ser -10	Pro	Ala	Gly	Pro	Arg -5	Tyr	Ser	Pro	Lys -1
Ser 1	Val	Ile	Gly	Ser 5	Asp	Glu	Arg	Thr	Arg 10	Val	Thr	Asn	Thr	Thr 15	Ala
Tyr	Pro	Tyr	Arg 20	Ala	Ile	Val	His	Ile 25	Ser	Ser	Ser	Ile	Gly 30	Ser	Суз
Thr	Gly	Ser 35	Leu	Ile	Gly	Pro	Lys 40	Thr	Val	Ala	Thr	Ala 45	Gly	His	Суз
Ile	Туr 50	Asp	Thr	Ala	Ser	Gly 55	Ser	Phe	Ala	Gly	Thr 60	Ala	Thr	Val	Ser
Pro 65	Gly	Arg	Asn	Gly	Ser 70	Thr	Tyr	Pro	Туr	Gly 75	Ser	Val	Thr	Ser	Thr 80
Arg	Tyr	Phe	Ile	Pro 85	Ser	Gly	Tyr	Arg	Ser 90	Gly	Asn	Ser	Asn	Tyr 95	Asp
Tyr	Gly	Ala	Ile 100	Glu	Leu	Ser	Gln	Pro 105	Ile	Gly	Asn	Thr	Val 110	Gly	Туг
Phe	Gly	Туг 115	Ser	Tyr	Thr	Thr	Ser 120	Ser	Leu	Val	Gly	Ser 125	Ser	Val	Thr
Ile	Ile 130	Gly	Tyr	Pro	Gly	Asp 135	Lys	Thr	Ser	Gly	Thr 140	Gln	Trp	Gln	Met
Ser 145	_	Asn	Ile	Ala	Val 150	Ser	Glu	Thr	Tyr	Lys 155	Leu	Gln	Tyr	Ala	11e
Asp	Thr	Tyr	Gly	Gly 165	Gln	Ser	Gly	Ser	Pro 170	Val	Tyr	Glu	Ala	Ser 175	Ser
Ser	Arg	Thr	Asn 180	Cys	Ser	Gly	Pro	Cys 185	Ser	Leu	Ala	Val	His 190	Thr	Asn

Gly Val Tyr Gly Gly Ser Ser Tyr Asn Arg Gly Thr Arg Ile Thr Lys

195 200 205

Glu Val Phe Asp Asn Leu Thr Asn Trp Lys Asn Ser Ala Gln 210 215 220

<210> 11

<211> 906

<212> DNA

<213> Bacillus pumilus JA96

<220>

<221> CDS

<222> (1)..(906)

<220>

<221> mat_peptide

<222> (262)..(906)

<220>

<221> sig_peptide

<222> (1)..(75)

<223> pro-peptide (76) ... (261)

<400> 11

atg aaa aag gtg aaa aaa tta atc cct tct cta ctc gtt ttt ggt gct 48
Met Lys Lys Val Lys Lys Leu Ile Pro Ser Leu Leu Val Phe Gly Ala
-85 -80 -75

tta agt gtg cct agt ttt gcc cat gca gca tct gat tca gta ctt acg 96
Leu Ser Val Pro Ser Phe Ala His Ala Ala Ser Asp Ser Val Leu Thr
-70 -65 -60

tet gat tat gac atg gtg act tet gac gga aag gtg att tet tea get

Ser Asp Tyr Asp Met Val Thr Ser Asp Gly Lys Val Ile Ser Ser Ala

-55

-40

gac ttc cac aac gat atg aaa acc ccc tca tcc ttt gac aaa gtg gat 192
Asp Phe His Asn Asp Met Lys Thr Pro Ser Ser Phe Asp Lys Val Asp
-35 -30 -25

gat ctc tct tct act att ggc gaa aaa gta aaa cca ctc aca aca tat 240
Asp Leu Ser Ser Thr Ile Gly Glu Lys Val Lys Pro Leu Thr Thr Tyr
-20 - -15 -10

tta aaa gac ttt caa aca aaa gta gtc att gga gac gat ggt aga aca 288 Leu Lys Asp Phe Gln Thr Lys Val Val Ile Gly Asp Asp Gly Arg Thr

		-5				-1					•					
aaa	ata	aco	aat	aca	aσa	σta	σca	ccc	tat	aat	tct	att	gct	tat	att	336
												Ile				
10	vui	••••			15	,,,			-,-	20					25	
aca	ttt	ggt	gga	tct	agc	tgc	act	gga	aca	ctc	att	gct	cca	aac	aaa	384
Thr	Phe	Gly	Gly	Ser 30	Ser	Суз	Thr	Gly	Thr	Leu	Ile	Ala	Pro	Asn 40	Lys	
				30					33					40		
ata	ttg	aca	aac	gga	cac	tgc	gtc	tac	aat	aca	gcc	aca	aga	agt	tat	432
Ile	Leu	Thr	Asn	Gly	His	Суз	Val	Tyr	Asn	Thr	Ala	Thr	Arg	Ser	Tyr	
			45					50					55			
												agc				480
Ser	Ala	Lys	Gly	Ser	Val	Tyr	Pro	Gly	Met	Asn	Asp	Ser	Thr	Ala	Val	
		60					65					70				
												agc				528
Asn	Gly	Ser	Ala	Asn	Met	Thr	Glu	Phe	Tyr	Val	Pro	Ser	Gly	Tyr	Ile	
	75					80					85					
												aaa				576
Asn	Thr	Gly	Ala	Ser	Gln	Tyr	Asp	Phe	Ala	Val	Ile	Lys	Thr	Asp		
90					95					100					105	
aac	att	gga	aat	acg	gtc	ggc	tat	cgc	tct	att	cgt	caa	gtg	aca	aat	624
Asn	Ile	Gly	Asn	Thr	Val	Gly	Tyr	Arg	Ser	Ile	Arg	Gln	Val	Thr	Asn	
				110					115					120		
cta	aca	ggt	aca	acg	att	aaa	att	tct	gga	tat	cca	ggt	gat	aaa	atg	672
Leu	Thr	Gly	Thr	Thr	Ile	Lys	Ile	Ser	Gly	Tyr	Pro	Gly	Asp	Lys	Met	
			125					130					135			
												ggt				720
Arg	Ser	Thr	Gly	Lys	Val	Ser	Gln	Trp	Glu	Met	Ser	Gly	Pro	Val	Thr	
		140					145					150				
aga	gaa	gat	acg	aat	ctc	gca	tac	tat	acg	atc	gat	aca	ttt	agc	gga	768
Arg	Glu	Asp	Thr	Asn	Leu	Ala	Tyr	Tyr	Thr	Ile	Asp	Thr	Phe	Ser	Gly	
	155					160					165					
												atc				816
Asn	Ser	Gly	Ser	Ala	Met	Leu	Asp	Gln	Asn	Gln	Gln	Ile	Val	Gly	Val	
170					175					180				-	. 185	
cat	aat-	gcg	ggt	tat	tca	aat	gga	acg	atc	aac	ggt	gga	cca	_aaa	gcg	864
ui c	λen	Δla	Glv	ጥላተ	Ser	Asn	GIV	Thr	Ile	Asn	Glv	Glv	Pro	Lys	Ala	

190 195 200

act gct gcc ttt gtt gaa ttt atc aac tat gcg aag gcg caa 906
Thr Ala Ala Phe Val Glu Phe Ile Asn Tyr Ala Lys Ala Gln
205 210 215

<210> 12

<211> 302

<212> PRT

<213> Bacillus pumilus JA96

<400> 12

Met Lys Lys Val Lys Lys Leu Ile Pro Ser Leu Leu Val Phe Gly Ala -85 -80 -75

Leu Ser Val Pro Ser Phe Ala His Ala Ala Ser Asp Ser Val Leu Thr
-70 -65 -60

Ser Asp Tyr Asp Met Val Thr Ser Asp Gly Lys Val Ile Ser Ser Ala
-55 -50 -45 -40

Asp Phe His Asn Asp Met Lys Thr Pro Ser Ser Phe Asp Lys Val Asp
-35 -30 -25

Asp Leu Ser Ser Thr Ile Gly Glu Lys Val Lys Pro Leu Thr Thr Tyr
-20 -15 -10

Leu Lys Asp Phe Gln Thr Lys Val Val Ile Gly Asp Asp Gly Arg Thr
-5 -1 1 5

Lys Val Thr Asn Thr Arg Val Ala Pro Tyr Asn Ser Ile Ala Tyr Ile 10 15 20 25

Thr Phe Gly Gly Ser Ser Cys Thr Gly Thr Leu Ile Ala Pro Asn Lys
30 35 40

Ile Leu Thr Asn Gly His Cys Val Tyr Asn Thr Ala Thr Arg Ser Tyr
45 50 55

Ser Ala Lys Gly Ser Val Tyr Pro Gly Met Asn Asp Ser Thr Ala Val 60 65 70

Asn Gly Ser Ala Asn Met Thr Glu Phe Tyr Val Pro Ser Gly Tyr Ile
75 80 85

Asn Thr Gly Ala Ser Gln Tyr Asp Phe Ala_Val Ile Lys Thr Asp Thr 90 95 100 105

Asn Ile Gly Asn Thr Val Gly Tyr Arg Ser Ile Arg Gln Val Thr Asn 115 Leu Thr Gly Thr Thr Ile Lys Ile Ser Gly Tyr Pro Gly Asp Lys Met 130 Arg Ser Thr Gly Lys Val Ser Gln Trp Glu Met Ser Gly Pro Val Thr 150 140 145 Arg Glu Asp Thr Asn Leu Ala Tyr Tyr Thr Ile Asp Thr Phe Ser Gly 155 160 Asn Ser Gly Ser Ala Met Leu Asp Gln Asn Gln Gln Ile Val Gly Val 175 180 170 His Asn Ala Gly Tyr Ser Asn Gly Thr Ile Asn Gly Gly Pro Lys Ala 190 195 Thr Ala Ala Phe Val Glu Phe Ile Asn Tyr Ala Lys Ala Gln 215 205 210 <210> 13 <211> 939 <212> DNA <213> Bacillus subtilis IS75 <220> <221> CDS <222> (1)..(939) <220> <221> mat_peptide <222> (280)..(939) <220> <221> sig_peptide <222> (1)..(102) <223> pro-peptide (103) ... (279) <400> 13 atg aaa tta gtt cca aga ttc aga aaa caa tgg ttc gct tac tta acg Met Lys Leu Val Pro Arg Phe Arg Lys Gln Trp Phe Ala Tyr Leu Thr

gtt ttg tgt ttg gct ttg gca gca gcg gtt tct ttt ggc gta ccg gca 96

-85

-90

Val	Leu	Cys -75	Leu	Ala	Leu	Ala	Ala -70	Ala	Val	Ser	Phe	Gly -65	Val	Pro	Ala	
					ccg Pro											144
					aac Asn -40											192
					gga Gly											240
					aac Asn											288
					acc Thr											336
_	-				ctg Leu 25											384
					ttt Phe											432
					agc Ser											480
					aat Asn											528
	_	-			tcc Ser											576
	Tyr				gct Ala 105											624
att	ggc	- tga	tac	ggc	tac	cgg	act	aca	aac	agc	agc	agt	ccc	gtg	ggc	672

Val Gly Trp	Tyr Gly Tyr 120	Arg Thr Thr	Asn Ser Ser Se 125	er Pro Val Gly 130
			tgt gac aaa ac Cys Asp Lys Th	
			tcc gct gaa ac Ser Ala Glu Th	
			caa agc ggc to Gln Ser Gly Se 175	
_			gct att gcc at Ala Ile Ala I 190	
			agg gtg acg as Arg Val Thr As 205	
		gca aat caa Ala Asn Gln 220		939
<210> 14 <211> 313 <212> PRT <213> Bacil	lus subtilis	IS75		
<400> 14 Met Lys Leu	Val Pro Arg	Phe Arg Lys	Gln Trp Phe A	la Tyr Leu Thr -80
Val Leu Cys -75		Ala Ala Ala -70	Val Ser Phe G	ly Val Pro Ala 55
Lys Ala Ala -60	Glu Asn Pro	Gln Thr Ser	Val Ser Asn T	nr Gly Lys Glu _.
Ala Asp Ala -45	Thr Lys Asn		Lys Ala Asp G	ln Val Ser Ala -30
Pro Tyr Glu	Gly Thr Gly	Lys Thr Ser	Lys Ser Leu T	yr Gly Gly Gln -15

Thr	Glu	Leu	-10	Lys	Asn	11e	GIn	-5	Leu	GIn	Pro	-1	ser 1	11e	iie
Gly	Thr 5	Asp	Glu	Arg	Thr	Arg 10	Ile	Ser	Ser	Thr	Thr 15	Ser	Phe	Pro	Tyr
Arg 20	Ala	Thr	Val	Gln	Leu 25	Ser	Ile	Lys	Tyr	Pro 30	Asn	Thr	Ser	Ser	Thr 35
Tyr	Gly	Cys	Thr	Gly 40	Phe	Leu	Val	Asn	Pro 45	Asn	Thr	Val	Val	Thr 50	Ala
Gly	His	Cys	Val 55	Tyr	Ser	Gln	Asp	His 60	Gly	Trp	Ala	Ser	Thr 65	Ile	Thr
Ala	Ala	Pro 70	Gly	Arg	Asn	Gly	Ser 75	Ser	Tyr	Pro	Tyr	Gly 80	Thr	Tyr	Ser
Gly	Thr 85	Met	Phe	Tyr	Ser	Val 90	Lys	Gly	Trp	Thr	Glu 95	Ser	Lys	Asp	Thr
Asn 100	Tyr	Asp	Tyr	Gly	Ala 105	Ile	Lys	Leu	Asn	Gly 110	Ser	Pro	Gly	Asn	Thr 115
Val	Gly	Trp	Tyr	Gly 120	Tyr	Arg	Thr	Thr	Asn 125	Ser	Ser	Ser	Pro	Val 130	Gly
Leu	Ser	Ser	Ser 135	Val	Thr	Gly	Phe	Pro 140	Суз	Asp	Lys	Thr	Phe 145	Gly	Thr
Met	Trp	Ser 150	Asp	Thr	Lys	Pro	Ile 155	Arg	Ser	Ala	Glu	Thr 160	Tyr	Lys	Leu
Thr	Tyr 165	Thr	Thr	Asp		Tyr 170	Gly	Cys	Gln	Ser	Gly 175	Ser	Pro	Val	Tyr
Arg 180	Asn	Tyr	Ser	qaA	Thr 185	Gly	Gln	Thr	Ala	Ile 190	Ala	Ile	His	Thr	Asn 195
Gly	Gly	Ser	Ser	туr 200	Asn	Leu	Gly	Thr	Arg 205	Val	Thr	Asn	Asp	Val 210	Phe
Asn	Asn	Ile	Gln 215	Tyr	Trp	Ala	Asn -	Gln 220							

```
<210> 15
<211> 4
<212> PRT
<213> Artificial Sequence
<220>
<223> Variation
<400> 15
Glu Lys Ala Ser
 1
<210> 16
<211> 6
<212> PRT
<213> Artificial Sequence
<220>
<223> Variation
<400> 16
Ser Glu Lys Ala Ser Thr
 1
<210> 17
<211> 4
<212> PRT
<213> Artificial Sequence
<220>
<223> Variation
<400> 17
Ser Glu Glu Thr
 1
<210> 18
<211> 38
<212> DNA
<213> Artificial Sequence
<220>
<223> Primer
<221> misc_feature
<222> (1)...(38)
\langle 223 \rangle n = A,T,C or G
```

```
<400> 18
                                                                        38
ggatggagaa gcggaaacac naaytaygay tayggngc
<210> 19
<211> 10
<212> PRT
<213> Artificial Sequence
<223> Variation
<400> 19
Gly Trp Arg Ser Gly Asn Tyr Asp Tyr Gly
<210> 20
<211> 28
<212> DNA
<213> Artificial Sequence
<220>
<223> Primer
<221> misc_feature
 <222> 12,15
<223> n = deoxyInosine
<221> misc_feature
 <222> 18,21 24
<223> n = A,T,C or G
<400> 20
                                                                        28
cccaagettg tngynacngc nggncayt
<210> 21
<211> 6
 <212> PRT
<213> Artificial Sequence
<220>
<223> Variation
<221> VARIANT
<222> 2<223> Xaa = Ala or Val
```

```
<400> 21
Val Xaa Thr Ala Gly His
<210> 22
<211> 36
<212> DNA
<213> Artificial Sequence
<220>
<223> Primer
<221> misc_feature
<222> (1)...(36)
<223> n = A,T,C or G
<400> 22
                                                                       36
gaataccggt gaaccgcttt gncmnccrta ngtrtc
<210> 23
<211> 12
<212> PRT
<213> Artificial Sequence
<220>
<223> Variation
<221> VARIANT
<222> 5
<223> Xaa = Gly or Cys or Trp
<400> 23
Asp Thr Tyr Gly Xaa Gln Ser Gly Ser Pro Val Phe
<210> 24
<211> 26
<212> DNA
<213> Artificial Sequence
<220>
<223> Primer
<221> misc_feature
<222> (1)...(26)
<223> n = A,T,C or G
```

```
<400> 24
                                                                        26
gctctagagt ydatngcncc rtartc
<210> 25
<211> 6
<212> PRT
<213> Artificial Sequence
<220>
<223> Variation
<221> VARIANT
<222> 6
<223> Xaa = Glu or Lys
<400> 25
Asp Tyr Gly Ala Ile Xaa
<210> 26
<211> 143
<212> DNA
<213> Bacillus
<400> 26
                                                                        60
gcgtctatga cacggcaagc cgatcattcg cgggaaccgc caccgtttcc ccgggacgaa
                                                                       120
acggttcagc ttacccttac ggatctgtta catcgacccg ctatttcatc ccgtcgggtt
                                                                       143
ggcagagcgg aaattccaat tat
<210> 27
<211> 48
<212> PRT
<213> Bacillus
<400> 27
Cys Val Tyr Asp Thr Ala Ser Arg Ser Phe Ala Gly Thr Ala Thr Val
                                                         15
                                    10
                 5
Ser Pro Gly Arg Asn Gly Ser Ala Tyr Pro Tyr Gly Ser Val Thr Ser
                                25
                                                     30
Thr Arg Tyr Phe Ile Pro Ser Gly Trp Gln Ser Gly Asn Ser Asn Tyr
                                                 45
        35
                            40
<210> 28
<211> 184
<212> DNA
<213> Bacillus
```

<400> 28 gatcgagete agecagecga teggeaatae egteggatat tteggatatt catacacege 60 ttcatcgctt gcaggagcag gcgtgaccat cagcggatat ccaggagaca aaacaacagg 120 180 cacccagtgg caaatgtccg gaacgatcgc tgtttcagaa acgtataaac tgcaatatgc 184 gatc <210> 29 <211> 61 <212> PRT <213> Bacillus <400> 29 Ile Glu Leu Ser Gln Pro Ile Gly Asn Thr Val Gly Tyr Phe Gly Tyr 15 5 Ser Tyr Thr Ala Ser Ser Leu Ala Gly Ala Gly Val Thr Ile Ser Gly 25 Tyr Pro Gly Asp Lys Thr Thr Gly Thr Gln Trp Gln Met Ser Gly Thr 40 Ile Ala Val Ser Glu Thr Tyr Lys Leu Gln Tyr Ala Ile 55 60 <210> 30 <211> 143 <212> DNA <213> Bacillus <400> 30 60 gcatttatga cacagcgagc gggtcattcg ccggaaccgc taccgtttct ccgggacgga acggttcaac atatccgtac ggatcagtta catcaacccg ctatttcatc ccgtcaggct 120 143 atcgaagcgg aaattcgaat tac <210> 31 <211> 48 <212> PRT <213> Bacillus Cys Ile Tyr Asp Thr Ala Ser Gly Ser Phe Ala Gly Thr Ala Thr Val 10 Ser Pro Gly Arg Asn Gly Ser Thr Tyr Pro Tyr Gly Ser Val Thr Ser 25 Thr Arg Tyr Phe Ile Pro Ser Gly Tyr Arg Ser Gly Asn Ser Asn Tyr 40 <210> 32 <211> 184 <212> DNA <213> Bacillus

<400> 32	
catagagete agecageega teggeaacae egtegggtat tteggatatt cetacacea	.c 60
ctcgtctctc gttgggtcaa gcgttaccat catcggatat ccaggcgaca aaacatcgg	g 120
cacccaatgg cagatgtccg gaaatatcgc cgtctcagaa acatataaac tgcaatatg	rc 180
gatc	184
<210> 33	
<211> 61	
<212> PRT	
<213> Bacillus	
<400> 33	
Ile Glu Leu Ser Gln Pro Ile Gly Asn Thr Val Gly Tyr Phe Gly Tyr	
1 5 10 15	
Ser Tyr Thr Thr Ser Ser Leu Val Gly Ser Ser Val Thr Ile Ile Gly	
20 25 30	
Tyr Pro Gly Asp Lys Thr Ser Gly Thr Gln Trp Gln Met Ser Gly Asn	
35 40 45	
Ile Ala Val Ser Glu Thr Tyr Lys Leu Gln Tyr Ala Ile	
50 55 60	
<210> 34	
<211> 19	
<212> DNA	
<213> Bacillus	
<400> 34	10
cgtaagggta agctgaacc	19
.010. 25	
<210> 35	
<211> 23	
<212> DNA <213> Bacillus	
<213> Bacillus	
<400> 35	
caggagacaa aacaacagca ggc	23
cayyayacaa aacaacayca yyc	-
<210> 36	
<211> 18	
<211> DNA	
<213> Bacillus	
<400> 36	
gtcccggaga aacggtag _	18
<210> 37	
<211> 20 "	

```
<212> DNA
<213> Bacillus
<400> 37
                                                                        20
caccacctcg tctctcgttg
<210> 38
<211> 29
<212> DNA
<213> Bacillus
<220>
<221> misc_feature
<222> 18,21
<223> Xaa = deoxyInosine
<221> misc_feature
<222> 27
<223> n = A, G, C or T
<400> 38
                                                                   29
gctctagacg tyttrtcncm nggrwancc
<210> 39
<211> 7
<212> PRT
<213> Artificial Sequence
<220>
<223> Variation
<221> VARIANT
<222> 2
<223> Xaa = is Tyr or Phe
<221> VARIANT
<222> 4
<223> Xaa = is Gly or Cys
<400> 39
Gly Xaa Pro Xaa Asp Lys Thr
```

<210> 40 <211> 29

<212> DNA _

<213> Artificial Sequence

```
<220>
 <223> Primer
 <221> misc_feature
 <222> 12,15,21
 <223> n = is deoxyInosine
 <400> 40
                                                                          29
 cccaagcttg tngtnathgg ngaygaygg
 <210> 41
 <211> 8
 <212> PRT
 <213> Artificial Sequence
 <220>
 <223> Variation
 <400> 41
 Val Val Ile Gly Gly Asp Asp Gly
  1
                  5
 <210> 42
 <211> 29
 <212> DNA
 <213> Artificial Sequence
 <220>
 <223> Primer
 <400> 42
                                                                          29
 gcacggaccg ttgcagttcg ttctggagc
 <210> 43<211> 33
 <212> DNA
 <213> Artificial Sequence
 <220>
 <223> Primer
 <400> 43
                                                                          33
 ccggcaaagt gaatgaaaca aaggaaaaag cgg
 <210> 44
 <211> 31
_ <212> DNA
```

<213> Artificial Sequence

<220>

<223> Primer

<221> misc_feature

<222> (1)...(31)

<223> n = A,T,C or G

<400> 44

atgcaccgga tggnnhatag gtccgaaaac c

31

<210> 45

<211> 21

<212> DNA

<213> Artificial Sequence

<220>

<223> Primer

<400> 45

ccctttaacc gcacagcgtt t

This Page is Inserted by IFW Indexing and Scanning Operations and is not part of the Official Record

BEST AVAILABLE IMAGES

Defective images within this document are accurate representations of the original documents submitted by the applicant.

Defects in the images include but are not limited to the items checked:
BLACK BORDERS
☐ IMAGE CUT OFF AT TOP, BOTTOM OR SIDES
☐ FADED TEXT OR DRAWING
☐ BLURRED OR ILLEGIBLE TEXT OR DRAWING
☐ SKEWED/SLANTED IMAGES
COLOR OR BLACK AND WHITE PHOTOGRAPHS
☐ GRAY SCALE DOCUMENTS
☐ LINES OR MARKS ON ORIGINAL DOCUMENT
REFERENCE(S) OR EXHIBIT(S) SUBMITTED ARE POOR QUALITY
OTHER:

IMAGES ARE BEST AVAILABLE COPY.

As rescanning these documents will not correct the image problems checked, please do not report these problems to the IFW Image Problem Mailbox.